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activation. The method comprises administering to the subject a therapeutically effective amount of a compound represented by structural formula (I) and physiologically acceptable salts thereof. Z is a substituted income form are to show the X is a covalent bord or of	d or un CO R _i c group	$-(CH_2)_n - X - N_R_b$ (1) substituted aromatic group. Y is a covalent bond, -O- or -CO n is an is an aliphatic or a substituted aliphatic group; R _b is an aliphatic group ; and, taken together with the nitrogen atom bonded to R _a and R _b , can

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CHEMOKINE RECEPTOR ANTAGONISTS AND METHODS OF USE THEREFOR

BACKGROUND OF THE INVENTION

Chemoattractant cytokines or chemokines are a family 5 of proinflammatory mediators that promote recruitment and activation of multiple lineages of leukocytes and lymphocytes. They can be released by many kinds of tissue cells after activation. Continuous release of chemokines at sites of inflammation mediates the ongoing migration of

- 10 effector cells in chronic inflammation. The chemokines characterized to date are related in primary structure. They share four conserved cysteines, which form disulfide bonds. Based upon this conserved cysteine motif, the family is divided into two main branches, designated as the C-X-C
- 15 chemokines (α-chemokines), and the C-C chemokines (β-chemokines), in which the first two conserved cysteines are separated by an intervening residue, or adjacent respectively (Baggiolini, M. and Dahinden, C. A., *Immunology Today*, 15:127-133 (1994)).
- 20 The C-X-C chemokines include a number of potent chemoattractants and activators of neutrophils, such as interleukin 8 (IL-8), PF4 and neutrophil-activating peptide-2 (NAP-2). The C-C chemokines include RANTES (Regulated on Activation, Normal T Expressed and 25 Secreted), the macrophage inflammatory proteins 1α and 1β

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(MIP-1 α and MIP-1 β), and human monocyte chemotatic proteins 1-3 (MCP-1, MCP-2, MCP-3), which have been characterized as chemoattractants and activators of monocytes or lymphocytes but do not appear to be chemoattractants for neutrophils. Chemokines, such as RANTES and MIP-1 α , have been implicated in a wide range of human acute and chronic inflammatory diseases including respiratory diseases, such as asthma and allergic disorders.

- The chemokine receptors are members of a superfamily 10 of G protein-coupled receptors (GPCR) which share structural features that reflect a common mechanism of action of signal transduction (Gerard, C. and Gerard, N.P., Annu Rev. Immunol., 12:775-808 (1994); Gerard, C. and Gerard, N. P., Curr. Opin. Immunol., 6:140-145 (1994)).
- 15 Conserved features include seven hydrophobic domains spanning the plasma membrane, which are connected by hydrophilic extracellular and intracellular loops. The majority of the primary sequence homology occurs in the hydrophobic transmembrane regions with the hydrophilic
- 20 regions being more diverse. The first receptor for the C-C chemokines that was cloned and expressed binds the chemokines MIP-1 α and RANTES. Accordingly, this MIP-1 α /RANTES receptor was designated C-C chemokine receptor 1 (also referred to as CCR-1; Neote, K., et al.,
- 25 Cell, 72:415-425 (1993); Horuk, R. et al., WO 94/11504, May 26, 1994; Gao, J.-I. et al., J. Exp. Med., 177:1421-1427 (1993)). Three new receptors have been characterized which bind and/or signal in response to RANTES: CCR3 mediates binding and signaling of chemokines including eotaxin,
- 30 RANTES, and MCP-3 (Ponath et al., J. Exp. Med., 183:2437 (1996)), CCR4 binds chemokines including RANTES, MIP-1α, and MCP-1 (Power, et al., J. Biol. Chem., 270:19495 (1995)), and CCR5 binds chemokines including MIP-1α,

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RANTES, and MIP-1 β (Samson, et al., Biochem. 35: 3362-3367 (1996)). RANTES is a chemotactic chemokine for a variety of cell types, including monocytes, eosinophils, and a subset of T-cells. The responses of these different cells

- 5 may not all be mediated by the same receptor, and it is possible that the receptors CCR1, CCR4 and CCR5 will show some selectivity in receptor distribution and function between leukocyte types, as has already been shown for CCR3 (Ponath *et al.*). In particular, the ability of RANTES to
- 10 induce the directed migration of monocytes and a memory population of circulating T-cells (Schall, T. et al., Nature, 347:669-71 (1990)) suggests this chemokine and its receptor(s) may play a critical role in chronic inflammatory diseases, since these diseases are 15 characterized by destructive infiltrates of T cells and
- 15 characterized by destructive infilt monocytes.

Many existing drugs have been developed as antagonists of the receptors for biogenic amines, for example, as antagonists of the dopamine and histamine receptors. No

- 20 successful antagonists have yet been developed to the receptors for the larger proteins such as chemokines and C5a. Small molecule antagonists of the interaction between C-C chemokine receptors and their ligands, including RANTES and MIP-1α, would provide compounds useful for inhibiting 25 harmful inflammatory processes "triggered" by receptor
- 25 harmful inflammatory processes "triggered" by receptor ligand interaction, as well as valuable tools for the investigation of receptor-ligand interactions.

SUMMARY OF THE INVENTION

30 It has now been found that a number of small organic molecules are antagonists of chemokine receptor function and can inhibit leukocyte activation and/or recruitment. An antagonist of chemokine receptor function is a molecule which can inhibit the binding of one or more chemokines,



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including C-C chemokines such as RANTES and MIP-1 α , to one or more chemokine receptors on leukocytes and/or other cell types. As a consequence, processes and cellular responses mediated by chemokine receptors can be inhibited with these

- 5 small organic molecules. Based on this discovery, a method of treating a subject with a disease associated with aberrant leukocyte recruitment and/or activation is disclosed. The method comprises administering to the subject a therapeutically effective amount of a compound or
- 10 small organic molecule which is an antagonist of chemokine receptor function. Compounds or small organic molecules which have been identified as antagonists of chemokine receptor function are discussed in detail hereinbelow, and can be used for the manufacture of a medicament for
- 15 treating or for preventing a disease associated with aberrant leukocyte recruitment and/or activation. The invention also relates to the disclosed compounds and small organic molecules and their use in treating or preventing a disease associated with aberrant leukocytes recruitment
- 20 and/or activation. The invention also includes pharmaceutical compositions comprising one or more of the compounds or small organic molecules which have been identified herein as antagonists of chemokine function and a suitable pharmaceutical carrier. The invention further
- 25 relates to novel compounds which can be used to treat an individual with a disease associated with aberrant leukocyte recruitment and/or activation.

BRIEF DESCRIPTION OF THE FIGURES

Figures 1A and 1B are histograms illustrating the inhibition by varying concentrations of LS370 and LS374 (also referred to herein as "L-370" and "L-374", respectively) in the chemotaxis of fresh peripheral blood mononuclear cells (PBMC) in response to RANTES or MIP-1α.

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DETAILED DESCRIPTION OF THE INVENTION

The present invention relates to small molecule compounds which are antagonists of chemokine receptor function. Accordingly, processes or cellular responses mediated by the binding of a chemokine to a receptor can be inhibited (reduced or prevented, in whole or in part), including leukocyte migration, integrin activation, transient increases in the concentration of intracellular free calcium [Ca^{**}]_i, and/or granule release of proinflammatory mediators.

The invention further relates to a method of treatment, including prophylactic and therapeutic treatments, of a disease associated with aberrant leukocyte recruitment and/or activation, including chronic

- 15 inflammatory disorders characterized by the presence of RANTES and/or MIP-1α responsive T cells, monocytes and/or eosinophils, including but not limited to diseases such as arthritis, psoriasis, multiple sclerosis, inflammatory bowel diseases such as ulcerative colitis and Crohn's
- 20 disease, as well as allergies and asthma. Other diseases associated with aberrant leukocyte recruitment and/or activation which can be treated (including prophylactic treatments) with the methods disclosed herein are inflammatory diseases associated with Human
- 25 Immunodeficiency Virus (HIV) infection, e.g., AIDS associated encephalitis, AIDS related maculopapular skin eruption, AIDS related interstitial pneumonia, AIDS related enteropathy, AIDS related periportal hepatic inflammation and AIDS related glomerulo nephritis. The method
- 30 comprises administering to a subject a therapeutically effective amount of a compound (i.e., one or more compounds) which inhibits chemokine receptor function, inhibits the binding of a chemokine to leukocytes and/or other cell types, and/or which inhibits leukocyte migration 35 to, and/or activation at, sites of inflammation. According

to the method, chemokine-mediated chemotaxis and/or activation of pro-inflammatory cells bearing receptors for chemokines can be inhibited. As used herein, "proinflammatory cells" includes but is not limited to

5 leukocytes, since chemokine receptors may be expressed on other cell types, such as neurons and epithelial cells.

In one embodiment of the present invention, the antagonist of chemokine receptor function is represented by Structural Formula (I):

$$Z \xrightarrow{\qquad } Y \xrightarrow{\qquad } (CH_2) \xrightarrow{n} X \xrightarrow{\qquad } N \xrightarrow{\qquad } R_a$$

10

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(I)

Z is a substituted or unsubstituted aromatic group.

Y is a covalent bond, -O- or -CO-.

n is an integer from one to about five. n is 15 preferably one, two, or three.

X is a covalent bond or -CO-.

 R_a and R_c , taken together with the nitrogen atom to which they are bonded, form a substituted or unsubstituted non-aromatic heterocyclic ring. For example, R_a and R_b , together with the nitrogen atom to which they are bonded, form a four, five, six, seven or eight-membered nitrogencontaining non-aromatic ring. Alternatively R_a is an

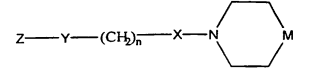
aliphatic or a substituted aliphatic group and R_b is an aliphatic group substituted with an aromatic group or

25 substituted aromatic group.

In a preferred embodiment, R_a and R_b , together with the nitrogen atom to which they are bonded, form a six-membered nitrogen-containing non-aromatic ring. For example, the

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six-membered, nitrogen-containing non-aromatic ring can be chosen such that the antagonist of chemokine receptor function is represented by Structural Formula (II):



(II)

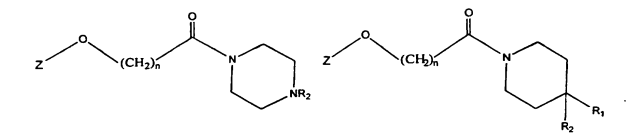
Z, Y, X and n are as described in Structural Formula (I).

M is >NR₂, >CR₁R₂, -O-, -S- or -CO-. M is preferably 10 >NR₂ or >CR₁R₂.

 R_1 is -H, -OH, an aliphatic group, -O-(aliphatic group), -SH or -S-(aliphatic group). Preferably, R_1 is -H or -OH.

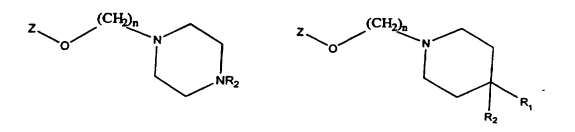
R₂ is an aliphatic group, a substituted aliphatic
15 group, an aromatic group, a substituted aromatic group, a benzylic group, a substituted benzylic group, a non-aromatic heterocyclic group or a substituted non-aromatic heterocyclic group.

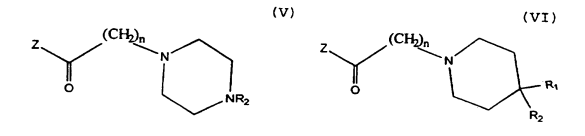
When M is $>NR_2$ or $>CR_1R_2$, the antagonist of chemokine 20 receptor function is preferably a compound represented by Structural Formulas (III) through (VIII):



(IV)

.





(VII)

(VIII)

In Structural Formulas (III) and (IV), n is preferably one, two or three, more preferably one. When n is one and R_1 is -H or -OH, R_2 is preferably a C_1 to about a C_4 alkyl group substituted with an aromatic or substituted aromatic 10 group.

In Structural Formulas (V) and (VI), n is preferably. . one, two or three, more preferably two or three. When n is two or three and R_1 is -H or -OH, R_2 is preferably an aliphatic or substituted aliphatic group, preferably an

15 alkyl group substituted with a hydroxyl, alkoxy, thiol, or alkylthiol group.

In Structural Formulas (VII) and (VIII), n is
preferably one, two or three, more preferably three. When
n is three and R₁ is -H or -OH, R₂ is preferably an
20 aromatic group, a substituted aromatic group or an

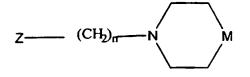
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aliphatic group substituted with an aromatic or substituted aromatic group.

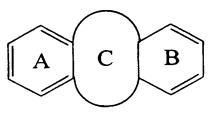
In another preferred embodiment, -X- and -Y- in Structural Formula (II) are each a covalent bond and the antagonist of chemokine receptor function is a compound represented by Structural Formula (IX):



(IX)

Z, n and M are as described above for Structural Formula

10 (II). Preferably, Z is a tricyclic ring system comprising two carbocyclic aromatic groups fused to a seven or eight membered cycloalkyl group or to a non-aromatic heterocyclic ring. In one example, Z is represented by Structural Formula (IXa):



(IXa)

The phenyl rings in Structural Formula (IXa), labeled with an "A" and "B", are referred to herein as "Ring A" and "Ring B", respectively. The central ring, labeled with a "C", is referred to as "Ring C" and can be, for example a seven or eight membered non-aromatic carbocyclic ring (e.g., a cycloheptane or cyclooctane ring) or a nonaromatic heterocyclic ring. When Ring C is a non-aromatic

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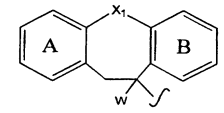
heterocyclic ring, it can contain one or two heteroatoms such as nitrogen, sulfur or oxygen. When Z is represented by Structural Formula (IXa), the tricyclic ring system is connected to the alkylene group in Structural Formula (IX)

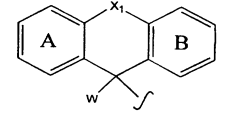
- 5 by a single covalent bond between the alkylene group and a ring atom in Ring C which is not also in Ring A or Ring B. Ring A and/or Ring B can be unsubstituted. Alternatively, Ring A and/or Ring B can have one or more substituents. Suitable substituents are as described
- 10 hereinbelow for substituted aromatic groups.

In addition, Ring C optionally contains one or more additional substituents, for example, R_3 and R_4 . When Ring C is a non-aromatic carbocyclic ring, substituents such as R_3 and R_4 are as described hereinbelow for substituted

- 15 aliphatic rings. When Ring C contains one or more heteroatoms, substituents such as R_3 and R_4 are as described below for non-aromatic heterocyclic rings. Preferably, R_3 is -H and R_4 is -H or an electron withdrawing group. Suitable electron withdrawing groups
- 20 include -CN, alkylsulfonyl, carboxamido, carboxylic alkyl esters, -NO₂ and halogens (e.g., -Br and -Cl).

More preferably, Z in Structural Formula (IX) is represented by Structural Formulas (X) and (XI):





25

(X)

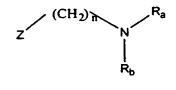
(XI)

 X_1 is a chemical bond, -S-, -CH₂- or -CH₂S-. Preferably, X_1 is -S- in Structural Formula (X) and -CH₂Sin Structural Formula (XI).

W is -H or an electron withdrawing group, as described above for Structural Formula (IXa). A preferred electron withdrawing group is -CN. Ring A and Ring B are as described above in Structural Formula (IXa).

When X_1 in Structural Formula (X) is -S- or when X_1 in Structural Formula (XI) is -CH₂S-, M is preferably >NR₂ or >CR₁R₂. When M is >NR₂ or >CR₁R₂, W is preferably -CN and n is preferably two, three or four, more preferably three. R₁ is preferably -H or -OH.

In another preferred embodiment, R_a is an aliphatic or a substituted aliphatic group and R_b is an aliphatic group 15 substituted with an aromatic group or substituted aromatic group. As a consequence, the antagonist of chemokine receptor function is a compound represented by Structural Formula (XII):



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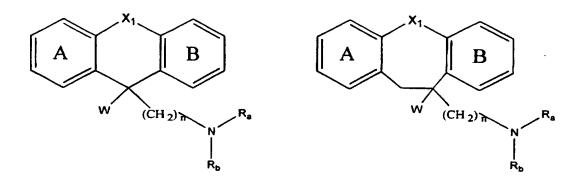
(XII)

Preferably, n is an integer from about two to about four; R_a is a C₁ to about a C₄ substituted or unsubstituted alkyl group; and R_b is $-(CH_2)_m-R_{10}$, wherein m is an integer from about two to about four, and R₁₀ is an aromatic group.

25

- In yet another preferred embodiment, the antagonist of chemokine function is a compound represented by Structural Formula (I), wherein Z is represented by Structural Formulas (X) or (XI) and -X- and -Y- are each a covalent bond. In this instance the antagonist of chemokine
- 30 receptor function is a compound represented by Structural Formulas (XIII) or (XIV):

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(XIII)

(XIV)

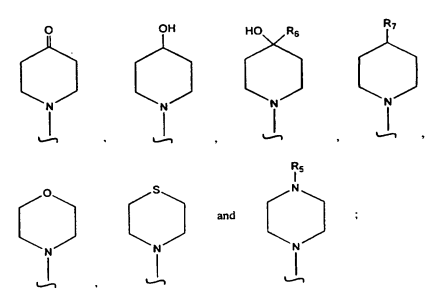
In Structural Formulas (XIII) and (XIV), X_1 , is as defined above for Structural Formulas (X) and (XI); n is an integer

5 from two to five; W is -H, -CN, alkylsulfonyl, carboxamido or carboxyalkyl;

In Structural Formulas (XIII) and (XIV), Ring A is substituted with R_8 and R_9 , wherein R_8 and R_9 are independently -H, a halogen, alkoxy or alkyl, or, taken

10 together with ring A, form a naphthyl group; and R_a and R_b are independently -H, alkyl, aralkyl or, taken together with the nitrogen atom bonded to R_a and R_b , form a non-aromatic heterocyclic ring represented by a structure selected from:

15



 $R_{\rm s}$ is -H, alkanoyl, aroyl, aralkoyl, alkyl, aralkyl or cycloalkanoyl.

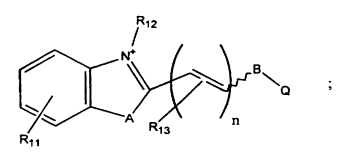
 R_6 is an aryl group.

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 R_{γ} is -H or a heterocylic ring.

In another embodiment of the present invention, the antagonist of chemokine receptor function is represented by Structural Formula (XVI):

. . . .



(XVI)

A is $>NR_{14}$, $-O_{-}$, $-S_{-}$, $-CH_{2}_{-}$, $-CH(R_{14})$ - or 5 $-C(R_{14}R_{15})$ -.

R₁₁ is -H, halogen, -CN, an aliphatic group, a substituted aliphatic group, an aromatic group, a substituted aromatic group, -OH, -O-(aliphatic group), -O-(substituted aliphatic group), -S-(aliphatic group), -S-

10 (substituted aliphatic group), -NO₂, -NH₂, -NH(aliphatic group), -NH(substituted aliphatic group), -N(aliphatic group)₂, -N(substituted aliphatic group)₂.

 R_{12} is an aromatic group or an aliphatic group.

Each R_{13} is independently chosen and is -H, an

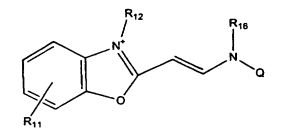
- 15 aliphatic group or substituted aliphatic group. Thus, if n is greater than one, the R_{13} attached to one double bond can be the same as or different from the R_{13} substituents attached to the other double bonds. Structural Formula (XVI) indicates that each R_{13} can be bonded to either
- 20 carbon atom in the double bond and that the stereochemistry of each double bond is independently selected and can be cis or trans.

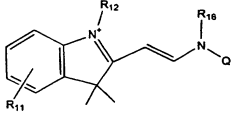
n is an integer from one to about four. B is $-N(R_{16})$ -, -S-, -O- or a covalent bond. R_{14} , R_{15} and R_{16} are independently an aliphatic group or a substituted aliphatic group and can be the same or different.

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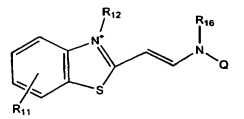
Q is an aliphatic group, a substituted aliphatic group, a benzylic group, a substituted benzylic group, an aromatic group, a substituted aromatic group, a nonaromatic heterocyclic group or a substituted non-aromatic heterocyclic group.

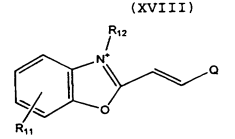
In a preferred embodiment, n is 1 and B and Q are as defined above. In this instance, A is preferably -O-, -Sor >C(CH₃)₂; B is -N(R₁₆)-, -S- or a covalent bond and R₁₃ is preferably -H or, when B is -S-, an aliphatic or substituted aliphatic group bonded to the same olefinic carbon atom as sulfur. As a consequence, the antagonist of chemokine receptor function is a compound represented by one of Structural Formulas (XVII) through (XXV):





(XVII)

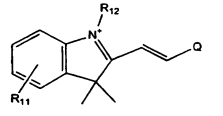


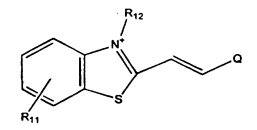


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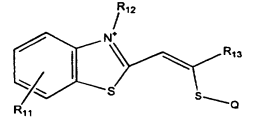
(XIX)

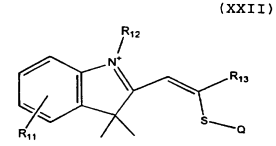
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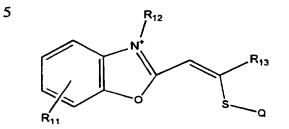
(XXI)







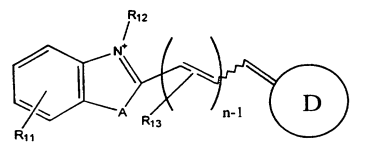




(XXV)

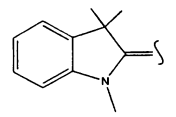
In Structures (XVII) through (XXV), R_{13} and R_{16} are preferably an aliphatic group.

Alternatively, in Structural Formula XVI, B, Q and the 10 terminal olefin carbon, taken together, form a non-aromatic heterocyclic ring. The antagonist of chemokine receptor function is then represented by Structural Formula (XXVI):

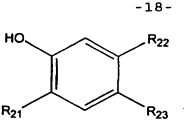


(XXVI)

- 5 R₁₁, R₁₂, R₁₃ and n are as described above for Structural Formula (XVI). Optionally, the non-aromatic heterocyclic ring in Structural Formula (XXVI), designated with a "D" and referred to herein as "Ring D", can be fused to an aromatic ring or substituted aromatic ring. The non-
- 10 aromatic heterocyclic ring can be substituted or unsubstituted. In one example, Ring D is represented by the following structural formula:



In another embodiment of the present invention, the 15 antagonist of chemokine receptor function is represented by Structural Formula (XXVII):



(XXVII)

R₂₁ is -OH, an aliphatic group, a substituted aliphatic
group, -O-(aliphatic group), -O-(substituted aliphatic group), -O-CO-(aliphatic group) or
-O-CO-(substituted aliphatic group). Preferably, R₂₁ is
-OH, CH₃CO-O- or an alkyl group substituted with CH₃NH-(e.g., an alkyl group substituted at the benzylic carbon
atom with methylamino methylene). Examples of R₂₁ include
-OH, CH₃CO-O- or -CH(-CH(CH₃)₂)(-CH₂NHCH₃).

 R_{22} and R_{23} are independently -H, an aliphatic group, a substituted aliphatic group, an aromatic group, a substituted aromatic group, -S-(aliphatic group), -S-

- 15 (substituted aliphatic group), -O-(aliphatic group), -O-(substituted aliphatic group), -(CH₂)_n-R₂₆, and, taken together, can be a -(CH₂)₂- to -(CH₂)₅- alkylene group or a -(CH₂)₂- to -(CH₂)₅- alkylene group substituted with one or more aliphatic groups, substituted aliphatic groups,
- 20 aromatic groups or substituted aromatic groups. Preferably, R_{22} is thioalkyl, alkyl or phenyl and R_{23} is -H, methyl or, taken together with R_{22} , a propylene group. The propylene group can be unsubstituted or substituted with one or more methyl or ethyl groups. Examples of R_{22}
- 25 include -SC₁H₁₅, methyl or phenyl. Examples of R₂₃ include -H, methyl or, taken together with R₂₂, a -CH₂CH₂C(CH₃)₂group.

 R_{26} is a substituted or unsubstituted aromatic group.

In one aspect, the antagonist of chemokine receptor function is a compound represented by Structural Formula (XXVII), wherein:

 R_{21} is -OH, an alkyl group, an alkoxy group, an acetoxy 5 group or an alkyl group substituted with -NR₂₄R₂₅;

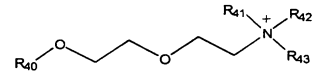
 R_{22} and R_{23} are independently an alkyl group, an aromatic group, an aralkyl group, and ethylene- R_6 or thioalkyl, and, taken together, form an alkylene group;

 R_{24} and R_{25} are independently an alkyl group, an aralkyl 10 group and an aryl group;

 R_{26} is a phenyl group substituted by R_{27} and $R_{28};$ and

 R_{27} and R_{28} are independently -H, -OH, alkoxy, or halogen.

In another embodiment of the present invention, the 15 antagonist of chemokine function is a compound represented by Structural Formula (XXVIII):



(XXVIII)

R₄₀ and R₄₃ are independently an aliphatic group, a substituted aliphatic group, a benzylic group, a substituted benzylic group, an aromatic group, a substituted aromatic group, a non-aromatic heterocyclic group or a substituted non-aromatic heterocyclic group.

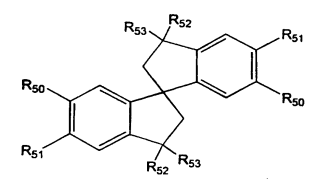
 R_{41} and R_{42} are independently an aliphatic group or a substituted aliphatic group. Preferably, R_{41} and R_{42} are each a methyl group.

In another embodiment of the present invention, the antagonist of chemokine receptor function is a compound represented by Structural Formula (XXIX):

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(XXIX)

R₅₀ and R₅₁ are independently -OH, a halogen, -O(aliphatic group), -O-(substituted aliphatic group), -O-CO(aliphatic group), -O-CO-(substituted aliphatic group),
-NH₂, -NH(aliphatic group), -NH(substituted aliphatic group), -N(aliphatic group)₂, -N(substituted aliphatic group)₂, -N(aliphatic group) or -S-(substituted aliphatic group)₂, -S-(aliphatic group) or -S-(substituted aliphatic group. Preferably, R₅₀ and R₅₁ are independently -OH, a
10 halogen, -O-(aliphatic group) or -O-(substituted aliphatic

group).

 R_{52} and R_{53} are independently -H, an aliphatic group, a substituted aliphatic group, a halogen, -NH₂, -NH(aliphatic group), -NH(substituted aliphatic group), -N(aliphatic

15 $group)_2$ or $-N(substituted aliphatic group)_2$. Preferably, R₅₂ and R₅₃ are independently an aliphatic group, a substituted aliphatic group or a halogen.

Also included in the present invention are physiologically acceptable salts of the compounds

20 represented by Structural Formulas (I) through (XXIX). Salts of compounds containing an amine or other basic group can be obtained, for example, by reacting with a suitable organic or inorganic acid, such as hydrogen chloride, hydrogen bromide, acetic acid, perchloric acid and the 5

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like. Compounds with a quaternary ammonium group also contain a counteranion such as chloride, bromide, iodide, acetate, perchlorate and the like.

As used herein, aliphatic groups include straight chained, branched or cyclic C_1-C_8 hydrocarbons which are completely saturated or which contain one or more units of unsaturation.

An "alkyl group" is a saturated aliphatic group, as defined above. The term "alkoxy" refers to an alkyl ether chain with an alkyl group. "Alkanoyl" refers to alkyl substituted carbonyl; "aralkanoyl" refers to phenyl-alkyl-CO- and "aroyl" refers to arylcarbonyl including benzoyl, naphthoyl and the like. The term "halogen" means fluoro, chloro, bromo and iodo. The term

- 15 "aryl", as opposed to the term "aromatic group", means phenyl. The term "substituted phenyl" means aryl substituted by alkyl, halogen, alkoxy, nitro, amino, acetamido, cyano and trifluoromethyl and naphthyl. "Aralkyl" means - (CH₂)_x-phenyl, wherein x is an integer from
- 20 one to four including benzyl. It is noted that the terms "aromatic group", "carbocylic aromatic group" and "heterocyclic aromatic group" are defined below and have different meanings from the term "aryl".

Aromatic groups include carbocyclic aromatic groups such as phenyl, 1-naphthyl, 2-naphthyl, 1-anthracyl and 2-

anthacyl, and heterocyclic aromatic groups such as Nimidazolyl, 2-imidazole, 2-thienyl, 3-thienyl, 2-furanyl, 3-furanyl, 2-pyridyl, 3-pyridyl, 4-pyridyl, 2-pyrimidy, 4pyrimidyl, 2-pyranyl, 3-pyranyl, 3-pyrazolyl, 4-pyrazolyl, 5-pyrazolyl, 2-pyrazinyl, 2-thiazole, 4-thiazole, 5-

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thiazole, 2-oxazolyl, 4-oxazolyl and 5-oxazolyl.

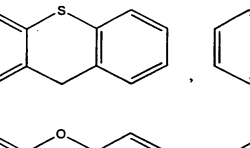
Aromatic groups also include fused polycyclic aromatic ring systems in which a carbocyclic aromatic ring or heteroaryl ring is fused to one or more other heteroaryl

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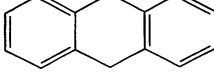
rings. Examples include 2-benzothienyl, 3-benzothienyl, 2-benzofuranyl, 3-benzofuranyl, 2-indolyl, 3-indolyl, 2quinolinyl, 3-quinolinyl, 2-benzothiazole, 2-benzooxazole, 2-benzimidazole, 2-quinolinyl, 3-quinolinyl, 1-

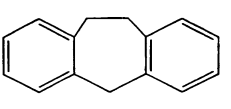
- 5 isoquinolinyl, 3-quinolinyl, 1-isoindolyl, 3-isoindolyl, and acridintyl. Also included within the scope of the term "aromatic group", as it is used herein, is a group in which one or more carbocyclic aromatic rings and/or heteroaromatic rings are fused to a cycloalkyl or non-
- 10 aromatic heterocyclic ring. Examples include decalin, phthalimido, benzodiazepines, benzooxazepines, benzooxazines, phenothiazines, and groups represented by the following structural formulas:

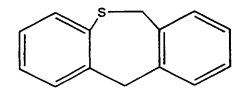
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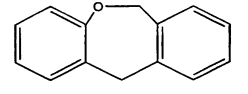


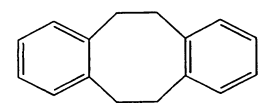
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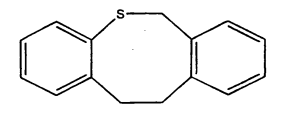


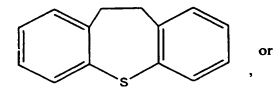


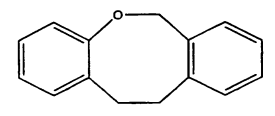












Non-aromatic heterocyclic rings are non-aromatic carbocyclic rings which include one or more heteroatoms such as nitrogen, oxygen or sulfur in the ring. The ring can be five, six, seven or eight-membered. Examples

- 5 include 2-tetrahydrofuranyl, 3-tetrahydrofuranyl, 2-tetrahyrothiophenyl, 3-tetrahyrothiophenyl, 2-morpholino, 3-morpholino, 4-morpholino, 2-thiomorpholino, 3-thiomorpholino, 4-thiomorpholino, 1-pyrrolidinyl, 2pyrrolidinyl, 3-pyrrolidinyl, 1-piperazinyl, 2-piperazinyl,
- 10 1-piperidinyl, 2-piperidinyl, 3-piperidinyl, 4-piperidinyl and 4-thiazolidinyl.

"Heterocyclic ring", as opposed to "heteroaryl group" and "non-aromatic heterocylic ring", is defined as imidazole, benzimidazole, pyridine, pyrimidine, thiazole,

15 benzothiazole, thienyl, benzothienyl. It is noted further the terms "heterocyclic aromatic group" and "non-aromatic heterocyclic ring" are defined above and have different meanings from the term "heterocyclic ring".

Suitable substituents on an alkyl, aliphatic, aromatic, non-aromatic heterocyclic ring or benzyl group include, for example, -OH, halogen (-Br, -Cl, -I and -F) -O(aliphatic, substituted aliphatic, benzyl, substituted benzyl, aromatic or substituted aromatic group), -CN, -NO₂,

-COOH, -NH₂, -NH(aliphatic group, substituted aliphatic,
benzyl, substituted benzyl, aromatic or substituted aromatic group), -N(aliphatic group, substituted aliphatic, benzyl, substituted benzyl, aromatic or substituted aromatic group)₂, -COO(aliphatic group, substituted aliphatic, benzyl, substituted benzyl, aromatic or

- 30 substituted aromatic group), -CONH₂, -CONH(aliphatic, substituted aliphatic group, benzyl, substituted benzyl, aromatic or substituted aromatic group)), -SH, -S(aliphatic, substituted aliphatic, benzyl, substituted benzyl, aromatic or substituted aromatic group) and -NH-
- 35 C(=NH)-NH₂. A substituted non-aromatic heterocyclic ring,

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benzylic group or aromatic group can also have an aliphatic or substituted aliphatic group as a substituent. A substituted alkyl or aliphatic group can also have a nonaromatic heterocyclic ring, benzyl, substituted benzyl,

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aromatic heterocyclic ring, benzyl, substituted benzyl, aromatic or substituted aromatic group as a substituent. A substituted non-aromatic heterocyclic ring can also have =0, =S, =NH or =N(aliphatic, aromatic or substituted aromatic group) as a substituent. A substituted aliphatic, substituted aromatic, substituted non-aromatic heterocyclic ring or substituted benzyl group can have more than one substituent.

In the structural formulas depicted herein, the single or double bond by which a chemical group or moiety is connected to the remainder of the molecule or compound is indicated by the following symbol:



For example, the corresponding symbol in Structural Formula (X) or (XI) indicates that the tricyclic ring system, which respresents Z in Structural Formula (IX), is connected to 20 the alkylene group in Structural Formula (IX) by a single covalent bond between the alkylene group and the ring carbon in Ring C which is bonded to W.

A "subject" is preferably a mammal, such as a human, but can also be an animal in need of veterinary treatment, 25 e.g., domestic animals (e.g., dogs, cats, and the like), farm animals (e.g., cows, sheep, pigs, horses, and the like) and laboratory animals (e.g., rats, mice, guinea pigs, and the like).

A "therapeutically effective amount" of a compound is 30 an amount which results in the inhibition of one or more processes mediated by the binding of a chemokine to a receptor in a subject with a disease associated with aberrant leukocyte recruitment and/or activation. Examples of such processes include leukocyte migration, integrin activation, transient increases in the concentration of intracellular free calcium $[Ca^{2+}]_i$ and granule release of

- 5 proinflammatory mediators. Alternatively, a "therapeutically effective amount" of a compound is a quantity sufficient to achieve a desired therapeutic and/or prophylactic effect, such as an amount which results in the prevention of or a decrease in the symptoms associated with
- 10 a disease associated with aberrant leukocyte recruitment and/or activation.

The amount of compound administered to the individual will depend on the type and severity of the disease and on the characteristics of the individual, such as general

- 15 health, age, sex, body weight and tolerance to drugs. It will also depend on the degree, severity and type of disease. The skilled artisan will be able to determine appropriate dosages depending on these and other factors. Typically, a therapeutically effective amount of the
- 20 compound can range from about 0.1 mg per day to about 100 mg per day for an adult. Preferably, the dosage ranges from about 1 mg per day to about 100 mg per day. An antagonist of chemokine receptor function can also be administered in combination with one or more additional
- 25 therapeutic agents, e.g. theophylline, β-adrenergic bronchdilators, corticosteroids, antihistamines, antiallergic agents and the like.

The compound can be administered by any suitable route, including, for example, orally in capsules,

- 30 suspensions or tablets or by parenteral administration. Parenteral administration can include, for example, systemic administration, such as by intramuscular, intravenous, subcutaneous, or intraperitoneal injection. The compound can also be administered orally (e.g.,
- 35 dietary), topically, by inhalation (e.g., intrabronchial,

intranasal, oral inhalation or intranasal drops), or rectally, depending on the disease or condition to be treated. Oral or parenteral administration are preferred modes of administration.

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The compound can be administered to the individual in conjunction with an acceptable pharmaceutical carrier as part of a pharmaceutical composition for treatment of HIV infection, inflammatory disease, or the other diseases discussed above. Formulation of a compound to be

- 10 administered will vary according to the route of administration selected (e.g., solution, emulsion, capsule). Suitable pharmaceutical carriers may contain inert ingredients which do not interact with the compound. Standard pharmaceutical formulation techniques can be
- 15 employed, such as those described in Remington's Pharmaceutical Sciences, Mack Publishing Company, Easton, PA. Suitable pharmaceutical carriers for parenteral administration include, for example, sterile water, physiological saline, bacteriostatic saline (saline)
- 20 containing about 0.9% mg/ml benzyl alcohol), phosphatebuffered saline, Hank's solution, Ringer's-lactate and the like. Methods for encapsulating compositions (such as in a coating of hard gelatin or cyclodextran) are known in the art (Baker, et al., "Controlled Release of Biological
- 25 Active Agents", John Wiley and Sons, 1986). The activity of compounds of the present invention can be assessed using suitable assays, such as receptor binding assays and chemotaxis assays. For example, as described in Exemplification Section, small molecule antagonists of 30 RANTES and MIP-1α binding have been identified utilizing
- 30 RANTES and MIP-1α binding have been identified utilizing HL-60 (butyric acid differentiated) cells which bind RANTES and chemotax in response to RANTES and MIP-1α as a model for leukocyte chemotaxis. Specifically, a high through-put receptor binding assay, which monitors ¹²⁵I-RANTES and



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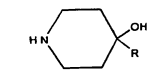
 125 I-MIP-1 α binding to HL-60 cell membranes, was used to identify small molecule antagonists which block binding and RANTES and MIP-1 α mediated HL-60, T-cell, peripheral blood mononuclear cell, and eosinophil chemotactic response.

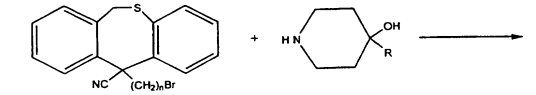
- 5 Compounds of the present invention can also be identified by virtue of their ability to inhibit the activation steps triggered by binding of a chemokine to its receptor, such as chemotaxis, integrin activation and/or granule mediator release.
- 10 The compounds represented by Structural Formula (IX), wherein Z is represented by Structural Formulas (IXa), (X) and (XI) and compounds represented by Structural Formulas (XIII) and (XIV) can be prepared according to methods described in Collect. Czech. Chem. Commun., 50(5):1089-96
- 15 (1985) (CA 104:33990) and Czech Patent CS 240698 B1 870601 (CA 109:92794). The teachings of these references and references cited therein are incorporated herein by reference. For example, these compounds can be prepared by the following reaction scheme:

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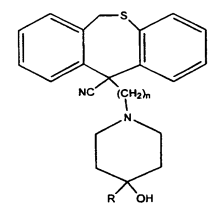








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Compounds represented by Structural Formula (V) and (VI), for example, the compounds designated in Table 1 as L-380 and Table 2 as L-372, can be prepared according to methods described in Collect. Czech. Chem. Commun.,

5 54(7):1966-1978 (1989), Czech Patent CS-268400 (1991) and WO 90/13539, the teachings of which are incorporated herein by reference.

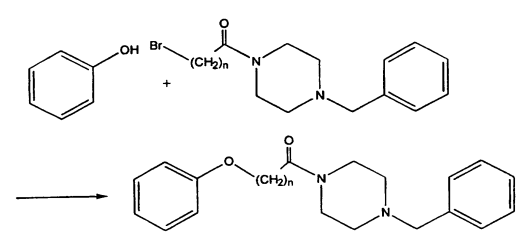
Compounds represented by Structural Formula (VII) and (VIII), for example, the compound designated as L-348 in

10 Table 2, can be prepared according to methods described in Synth. Commun. 25(2):177-82 (1995), Chem. Lett., (12):2295-8 (1994), Ther. Drug. Monit. 10(2):177-83 (1988), J. Med. Chem. 28(9):1319-24 (1985), U.S. Patent 4,086,234, U.S. Patent 4,012,514, U.S. Patent 3,936,468, U.S. Patent

15 3,922,266 and U.S. Patent 3,907,812, the teachings of which are incorporated herein by reference.

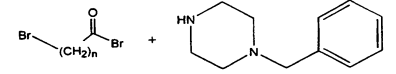
Compounds represented by Structural Formula (III) and (IV), for example the compound designates as L-377 in Table 2, can be prepared according to methods well known in the

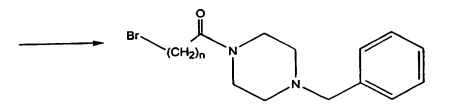
20 field of organic chemistry, for example, by reacting the sodium salt of a suitable phenol and a suitable alkylating agent. The phenol is preferably substituted with electron withdrawing groups (e.g., 3,4,5-trimethoxyphenol). This reaction is shown schematically below:



The phenol in the scheme above is preferably substituted with one or more electron withdrawing groups. The alkylating agent prepared, for example, by reacting a suitable bromo substituted acyl bromide (e.g., bromoacetyl

5 suitable bromo substituted acyl bromide (e.g., bromoacety bromide) with a suitable 1-substituted piperazine, for example, 1-benzylpiperazine, as shown below:





Compounds represented by Structural Formula (XII), for example, the compound designated L-347 in Table 1, can be prepared according to methods described in WO 97/11938, WO 97/09983, WO 96/40097, WO 96/39407 and EP 694543,

5 the teachings of which are incorporated herein by reference.

Compounds represented by Structural Formula (XVIII) and (XXI), for example the compound designated L-344 in Table 2, can be prepared, for example, by reacting a 1,3,3-

- 10 trialkylindolinium anion with a suitable alkylating agent according to methods described in European Patent 94 EP 0400348 and U.S. Patent No. US 5,258,274, the teachings of which are incorporated herein by reference. By replacing the 1,3,3-trialkylindolinium anion with an
- 15 appropriate 1-alkyl-benzoxazolinium anion or 1-alkylbenzothiazolinium anion, similar procedures can be used to prepare compound represented by Structural Formulas (XVII), (XIX), (XX) and (XXII) through (XXV) (e.g., compounds designated as L-459 and L-464 in Table II). These
- 20 procedures are also suitable for preparing compounds represented by Structural Formula (XXVI), for example, the compound designated L-342 in Table 2, by using an appropriate alkylating agent.

Compounds represented by Structural Formula (XXVII), 25 for example, the compound designated L-381 in Table 1, can be prepared according to methods described in EP 757982, EP 533056, EP 457701, EP 434093 and EP 332064, the teachings of which are incorporated herein by reference. Other compounds represented by Structural

30 Formula (XXVII), for example, the compound designated L-345 in Table 1, can be prepared according to methods described in Sb. Pr. Vyzk. Chem. Vyuziti Uhli, Dehtu Ropy 7:21-39 (1967), Z. Naturforsch. B: Anorg. Chem. Org. Chem 34B(4):624-32 (1979) and J. Med. Chem. 26(6):823-31 (1983),

the teachings of which are incorporated herein by reference. Yet other compounds represented by Structural Formula (XXVII), for example, the compound designated L-349 in Table 1, can be prepared according to methods described in EP 707007, WO 9501326, EP596692, EP 587050, EP 540165 and CA 2028031, the teachings of which are incorporated herein by reference.

Compounds represented by Structural Formula (XXVIII), for example, the compound designated L-339 in Table 1, can be prepared according to methods described in WO 94/26302, *Collect. Czech. Chem. Commun.* 53(7):1424-60 (1988), EP 226448, ES 540861 and Bull. Chem. Soc. Jap 44(6):1560-2, the teachings of which are incorporated herein by reference.

- 15 Compounds represented by Structural Formula (XXIX), for example, the compound designated L-319 in Table 1, can be prepared according to methods described in JP 09110771, Polym. Mater. Sci. Eng. 70:378-9 (1993), JP 03148232, JP 02286642, JP 03386641, JP 02248954, EP 342035, EP 307951,
- 20 Eur. Polym. J. 15(7):631-8 (1979), FR 2322161, Izv. Akad. Nauk. SSSR. Ser. Khim. (12):2808-10 (1973) and Tetrahedron Lett. (34):3707:10 (1968), the teachings of which are incorporated herein by reference.

The invention is illustrated by the following examples 25 which are not intended to be limiting in any way.

EXEMPLIFICATION

Human eosinophils were prepared by isolation from the blood of donor individuals with high levels of circulating blood eosinophils (5-17%) by combining density gradient centifugation and negative selection with anti-CD16 magnetic beads (Hansel, T.T. J. Immunol. Methods, 122:97-

103 (1989)). Briefly, the granulocyte fraction from the Percoll centrifugation was incubated with CD16 micro beads (miniMACS, separation unit) for 30 minutes. Cells were then passed through a MACS column (Miltenyi Biotec, Inc., Auburn, CA) and eosinophils were collected in the flow through. Eosinophil purity was >99% as determined by analysis of Diff-Quik (Baxter) stained cytocentrifugation preparations by light microscopy.

HL-60 Cells, obtained from the American Type Culture Collection, were resuspended at 0.5 million cells/ml in equal proportions of RPMI-1640 and M199 (Gibco) with 20% fetal calf serum (FCS). After, addition of n-butyric acid (Sigma Chemical Co.) to a final concentration of 0.4 mM, cells were incubated for 4 days at 37°C, 5%CO₂ before use

15 in either whole cell chemotaxis assays or preparation for use as membranes for receptor binding assays.

Membrane Preparations for Chemokine Binding and Binding Assays

Membranes were prepared from n-butyric acid-treated HL60

- 20 cells. Cells were harvested by centrifugation, washed twice with PBS (phosphate-buffered saline), and the cell pellets were frozen at -70 to -85°C. The frozen pellet was thawed in ice-cold lysis buffer consisting of 5 mM HEPES (N-2-hydroxyethylpiperazine-N'-2-ethane-sulfonic acid) pH
- 25 7.5, 2 mM EDTA (ethylenediaminetetraacetic acid), 5 μ g/ml each aprotinin, leupeptin, and chymostatin (protease inhibitors), and 100 μ g/ml PMSF (phenyl methane sulfonyl fluoride - also a protease inhibitor), at a concentration of 1 to 5 x 10' cells/ml. This procedure results in cell
- 30 lysis. The suspension was mixed well to resuspend all of the frozen cell pellet. Nuclei and cell debris were removed by centrifugation of 400 x g for 10 minutes at 4°C.

The supernatant was transferred to a fresh tube and the membrane fragments were collected by centrifugation at 25,000 x g for 30 minutes at 4°C. The supernatant was aspirated and the pellet was resuspended in freezing buffer consisting of 10 mM HEPES pH 7.5, 300 mM sucrose, $1\mu g/ml$ 5 each aprotinin, leupeptin, and chymostatin, and 10 μ g/ml PMSF (approximately 0.1 ml per each 10° cells). All clumps were resolved using a minihomogenizer, and the total protein concentration was determined using a protein assay kit (Bio-Rad, Hercules, CA, cat #500-0002). The membrane solution was then aliquoted and frozen at -70 to -85°C until needed.

Binding Assays utilized the membranes described above. Membrane protein (2 to 20 μ g total membrane protein) was

incubated with 0.1 to 0.2 nM $^{125}\text{I}\text{-labeled}$ RANTES or MIP-1 α 15 with or without unlabeled competitor (RANTES or MIP-1 α) or various concentrations of compounds. The binding reactions were performed in 60 to 100 μ l of a binding buffer consisting of 10 mM HEPES pH 7.2, 1 mM $CaCl_2$, 5 mM $MgCl_2$,

- and 0.5% BSA (bovine serum albumin), for 60 min at room 20 temperature. The binding reactions were terminated by harvesting the membranes by rapid filtration through glass fiber filters (GF/B or GF/C, Packard) which were presoaked in 0.3% polyethyleneimine. The filters were rinsed with
- approximately 600 μ l of binding buffer containing 0.5 M 25 NaCl, dried, and the amount of bound radioactivity was determined by scintillation counting in a Topcount betaplate counter.

Chemokines and Chemotaxis.

RANTES and MIP-1 α were purchased from Peprotech, Inc. 30 Leukocyte chemotaxis was assessed on eosinophils, peripheral blood mononuclear cells, or HL60 cells

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differentiated with butyric acid, using a modification of a transendothelial assay (Carr, M.W., et al. T.A., Proc. Natl Acad Sci, USA, <u>91</u>, 3652 (1994)). The endothelial cells used in this assay were the endothelial cell line, ECV 304,

- 5 obtained from the European collection of Animal Cell Cultures (Porton Downs, Salisbury, U.K.). Endothelial cells were cultured on 6.5 mm diameter Transwell culture inserts (Costar Corp., Cambridge,MA) with 3.0 μm pore size. Culture media for the ECV 304 cells consisted of M199+10%
- 10 FCS, L-glutamine, and antibiotics. The assay media consisted of equal parts RPMI 1640 and M199 with 0.5% BSA. Two hours before the assay, 2x10⁵ ECV 304 cells were plated onto each insert of the 24 well Transwell chemotaxis plate and incubated at 37°C. Chemotactic factors such as RANTES
- 15 or MIP-1 α (Peprotech) (diluted in assay medium) were added to the 24-well tissue culture plates in a final volume of 600 μ L. Endothelial-coated Transwells were inserted into each well and 10⁶ cells of the leukocyte type being studied were added to the top chamber in a final volume of 100 μ L of
- 20 assay medium. The plate was then incubated at 37°C in 5% $CO_2/95$ % air for 1-2h. The cells that had migrated to the bottom chamber were counted using flow cytometry. 500μ L of the cell suspension from the lower chamber was placed in a tube and relative counts were obtained for a set period of
- 25 time of 30 seconds. This counting method was found to be highly reproducible and enabled gating on the leukocytes and the exclusion of debris or other cells. Counts obtained by this method matched closely those obtained by counting with a microscope. Assays evaluating chemotaxis inhibitors
- 30 were performed in the same way as control experiments above, except that inhibitor solutions, in assay media containing up to 1% of DMSO cosolvent, were added to both the top and bottom chambers prior to addition of the cells. Inhibitor potency was determined by comparison of cell

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numbers migrated to the bottom chamber, with or without inhibitor. Control wells contained equivalent amounts of DMSO, but no inhibitor.

Ligand Binding Assay.

- 5 ¹²⁵I-RANTES and ¹²⁵I-MIP-1α were purchased from DuPont-NEN (Boston,MA) with a specific activity of 2,200 Ci/mM. Chemokine binding to the target cells, human eosinophils, was carried out using a modification of a method previously reported.(Van Riper, G.S.; J. Exp. Med. 177, 851-856
- 10 (1993)). Cells were washed once in PBS and resusupended in binding buffer (50mM HEPES, 1 mM CaCl₂, 5 mM MgCl₂, and 0.5% BSA) at a concentration of 1×10^7 / mL. Aliquots of 50 μ L (5x10⁵ cells) were dispensed into microfuge tubes, followed by the addition of cold and radiolabelled
- 15 chemokines. The final reaction volume was 200 μ L. Nonspecific binding was determined by incubating cells with radiolabeled chemokines in the presence of increasing amounts of (250-500 nM) of cold chemokine. After 60-min incubation, at room temperature, the cells were washed 3x
- 20 with 1 mL of binding buffer plus 0.5 M NaCl. Cell pellets were then counted. All experiments were carried out using duplicates and repeated at least three times. Curve fit was calculated by Kaleidagraph software (Synergy Software, Reading, PA). Inhibition of binding was assessed by the
- 25 addition of test inhibitor compound at concentrations of 100 μ M final concentration, and incubation for 30 min prior to addition of the chemokine as above.

Inhibition of Peripheral Blood Mononuclear Cell (PBMC) Chemotaxis By Compounds L-370 and L-374

30 Cells were incubated with the concentrations of compound indicated in Figures 1A and 1B for 20 minutes at room temperature and were placed in the upper wells of the

chemotaxis chambers. Migration in response to MCP-1, RANTES, or MIP-1 α was assessed as described above.

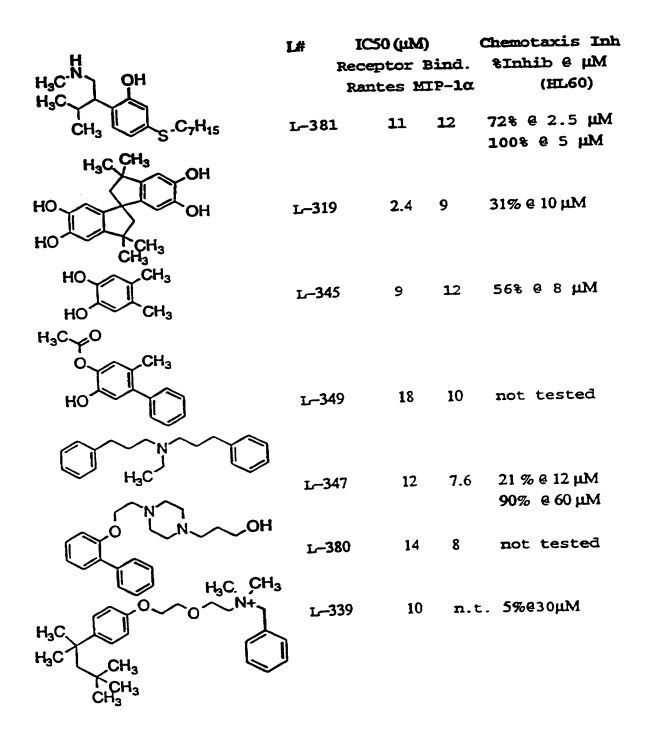
Figure 1A is an illustration of the total number of cells migrating in response to the chemokines with and without preincubation with different concentrations of L-370 or L-374. MCP-1 was used as a negative control to show the specificity of action of the compounds.

Figure 1B is an illustration of the results of the same experiments as in Figure 1A, expressed as percentage 10 inhibition, where the inhibition was calculated as cells migrated in the absence of compound/cells migrated in the presence of compound. 100% inhibition of migration occurred with 10.0 μ M and 1.0 μ M of L-370 and L-374, respectively.

- 15 The activities of other test compounds are reported in Tables 1-4 below as RBA, IC_{50} or the inhibitor concentration required for 50% inhibition in receptor binding assays using ¹²⁵I-RANTES or ¹²⁵MIP-1 α as ligand and HL60 cell membranes from cells differentiated by butyric 20 acid (which chemotax in response to RANTES in an almost
- identical way described for eosinophils).

Leukocyte chemotaxis inhibition is expressed as percent inhibition of RANTES-induced chemotaxis using the same HL60 cells (butyric acid diffentiated) at the indicated concentration (μ M) of compound.

Table 1



	-40-			
	Table	2		
	L#	RBA		Inhibition
	,		M)	HL60 Chemotaxis
			s MIP-1a	%Inhib'n, μM
С Н Он	l-377	2	0.6	66%810µМ
CH ₃ O CH ₃	1 33 9	10	23	not tested
	L-37 2	5 .5	17	89%е6µМ
F-{}-{~n}+	L-348	8	10	54%@4µM 103%@20µM
CH ₃ H ₃ C CH ₃ H ₃ C -N CH ₃ H ₄ C-N CH ₃	L-342	6	≡12	74%@2µM
	L-344	5	15	67%@2µM
H ₃ C, CH ₃ CH ₃	L~459	3	13	92%@15µM
	L 46 4	35	≡10	not tested

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-41-Table 3

NC OH	L#	RBA IC50 (µM)		Leukocyte Chemotaxis (HL60 Cells)
^N X		RANTES	MIP-1a	% Inhibition @ µM
	L-886	11.3	11.2	not tested
	L-804	>20	not tested	not tested
NC NC OH			0.36	81% @1µM
	L-374	0.2	0.20	
	L-370	7.3	11.7	59 %62µM 102%610 µM
	L-887	>40	not tested	not tested
	L-378	21	33	not tested
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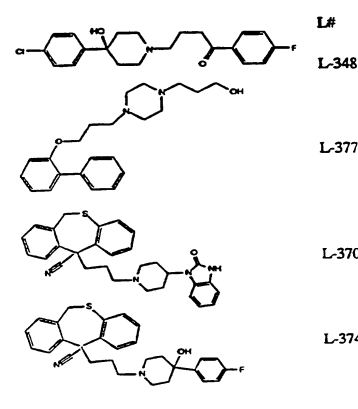
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-42-

Table 4

INHIBITION OF EOTAXIN-INDUCED EOSINOPHIL CHEMOTAXIS



17%/7μM
86% / 35 µM

L-377	100% f 3 µM
	100% / 6 µM

ł

L-370	26%/2μM
	40% / 10 µM

L-374 IC50 = $45.5 \,\mu\text{M}$

Equivalents

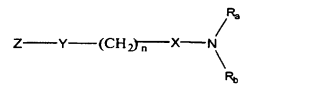
Those skilled in the art will be able to recognize, or be able to ascertain, using no more than routine experimentation, many equivalents to the specific

5 embodiments of the invention described herein. Such equivalents are intended to be encompassed by the following claims.

CLAIMS

What is claimed:

 A method of treating a subject with a disease associated with aberrant leukocyte recruitment and/or activation comprising administering to the individual a therapeutically effective amount of a compound represented by the following structural formula:



and physiologically acceptable salts thereof, 10 wherein:

Z is a substituted or unsubstituted aromatic group;

Y is a covalent bond, -O- or -CO-;

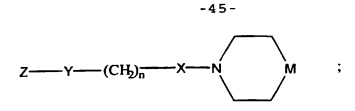
n is an integer from one to about five;

X is a covalent bond or -CO-; and

R_a is an aliphatic or a substituted aliphatic group; and

 R_b is an aliphatic group substituted with an aromatic group or substituted aromatic group; and wherein R_a and R_b , taken together with the nitrogen atom bonded to R_a and R_b , can form a substituted or unsubstituted non-aromatic heterocyclic ring.

2. The method of Claim 1 wherein the compound is represented by the following structural formula:



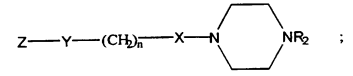
and physiologically acceptable salts thereof, wherein:

M is $>NR_2$, $>CR_1R_2$, -O-, -S- or -CO-;

R₁ is -H, -OH, an aliphatic group, -O-(aliphatic group), -SH or -S-(aliphatic group);

 R_2 is an aliphatic group, a substituted aliphatic group, an aromatic group, a substituted aromatic group, a benzylic group, a substituted benzylic group, a non-aromatic heterocyclic group or a substituted non-aromatic heterocyclic group.

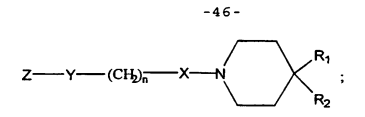
3. The method of Claim 2 wherein the compound is represented by the following structural formula:



and physiologically acceptable salts therof.

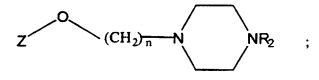
- The method of Claim 3 wherein -Y- is -O- and -X- is -CO-.
- 20 5. The method of Claim 4 wherein n is one and R_2 is a Cl to about a C4 alkyl group substituted with an aromatic or substituted aromatic group.
 - 6. The method of Claim 2 wherein the compound is represented by the following structural formula:

15



and physiologically acceptable salts thereof, wherein R_1 is -H or -OH.

- 7. The method of Claim 6 wherein -Y- is -O- and -X- is -CO-.
 - 8. The method of Claim 7 wherein n is one and R_2 is C1 to about a C4 alkyl group substituted with an aromatic or substituted aromatic group.
- 9. The method of Claim 3 wherein the compound is represented by the following structural formula:



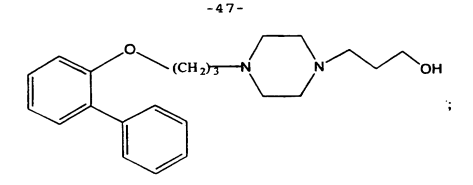
and physiologically acceptable salts thereof.

- 10. The method of Claim 9 wherein n is 2 or 3 and R_2 is an aliphatic or substituted aliphatic group.
- 15

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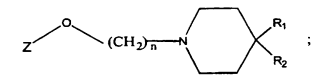
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11. The method of Claim 9 wherein the compound is represented by the following structural formula:



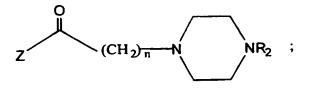
and physiologically acceptable salts thereof.

12. The method of Claim 6 wherein the compound is represented by the following structural formula:



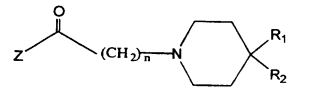
and physiologically acceptable salts thereof, wherein R_1 is -H or -OH.

- 13. The method of Claim 12 wherein n is two or three and R_2 is an aliphatic or substituted aliphatic group.
- 10 14. The method of Claim 3 wherein the compound is represented by the following structural formula:



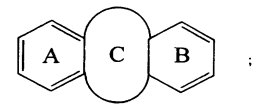
and physiologically acceptable salts thereof.

- 15. The method of Claim 14 wherein n is 3 and R_2 is an aromatic group, a substituted aromatic group or an aliphatic group substituted with an aromatic or substituted aromatic group.
- 5 16. The method of Claim 6 wherein the compound is represented by the following structural formula:



and physiologically acceptable salts thereof.

- 17. The method of Claim 16 wherein n is 3 and R₂ is an
 aromatic group, a substituted aromatic group or an aliphatic group substituted with an aromatic or substituted aromatic group.
 - 18. The method of Claim 2 wherein -X- and -Y- are each a covalent bond.
- 15 19. The method of Claim 18 wherein Z is represented by the following structural formula:

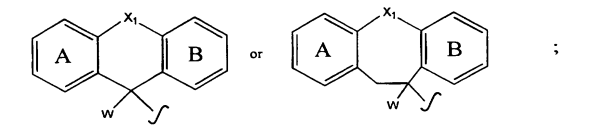


(IXa)

wherein:

Ring C is a substituted or unsubstituted C, or C_8 non-aromatic carbocyclic ring or a substituted or unsubstituted non-aromatic heterocyclic ring and is bonded to the alkylene group by a single covalent bond between the alkylene group and a ring atom in Ring C which is not also in Ring A or Ring B.

20. The method of Claim 18 wherein Z is represented by a structural formula selected from:



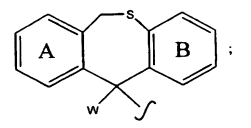
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wherein:

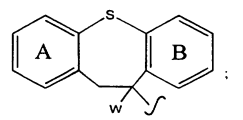
 X_1 is a chemical bond, -S-, -CH₂- or -CH₂S-; W is -H or an electron withdrawing group; and wherein ring A and ring B are substituted or estituted.

- 15 unsubstituted.
 - 21. The method of Claim 20 wherein Z is represented by the following structural formula:

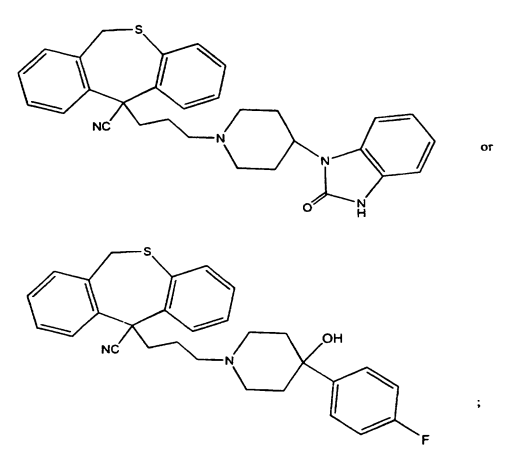


wherein Ring A and/or Ring B are substituted or unsubstituted.

- 22. The method of Claim 21 wherein M is $>NR_2$, $>C(OH)'R_2$ or $>CHR_2$.
- 5 23. The method of Claim 22 wherein n is three and W is -CN.
 - 24. The method of Claim 20 wherein Z is represented by the following structural formula:



- 10 wherein Ring A and/or Ring B are substituted or unsubstituted.
 - 25. The method of Claim 24 wherein M is $>C(OH)R_2$ or $>CHR_2$.
 - 26. The method of Claim 25 wherein W is -CN and n is three.
- 15 27. The method of Claim 1 wherein the compound is represented by a structural formula selected from:



and physiologically acceptable salts thereof.

28. The method of Claim 1 wherein: -X- and -Y- are each a covalent bond; Z is represented by a structural formula selected

from: $A \rightarrow B$ or $A \rightarrow B$;

5 wherein: X₁ is a chemical bond, -S-, -CH₂- or -CH₂S-; W is -H, -CN, alkylsulfonyl, carboxamido or carboxyalkyl;

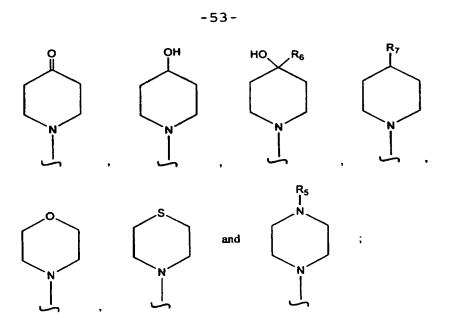
n is an integer from 2-5;

Ring A is substituted with R_8 and R_9 , wherein R_8 and R_9 are independently -H, a halogen, alkoxy or alkyl, or, taken together with ring A, form a naphthyl group;

 R_a and R_b are independently -H, alkyl, aralkyl or, taken together with the nitrogen atom bonded to R_a and R_b , form a non-aromatic heterocyclic ring represented by a structure selected from:

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and physiologically acceptable salts thereof, wherein:

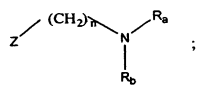
 R_5 is -H, alkanoyl, aroyl, aralkoyl, alkyl, 5 aralkyl or cycloalkyl;

 R_6 is an aryl group; and

 R_7 is -H or a heterocylic ring.

29. The method of Claim 1 wherein the compound is represented by the following structural formula:

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and physiologically acceptable salts thereof.

30. The method of Claim 29; wherein:

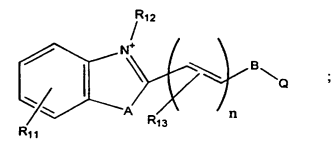
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n is an integer from about 2-4;
```

 R_a is a C1 to about C4 substituted or unsubstituted alkyl group; and

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 R_b is $-(CH_2)_m - R_{10}$ wherein m is an integer from about 2-4 and R_{10} is an aromatic group.

- 31. A method of treating a subject with a disease associated with aberrant leukocyte recruitment and/or activation comprising administering to the individual
- 10 a therapeutically effective amount of a compound represented by the following structural formula:



and physiologically acceptable salts thereof, wherein:

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A is $>NR_{14}$, $-O_{-}$, $-S_{-}$, $-CH_{2}_{-}$, $-CH(R_{14})$ - or $-C(R_{14}R_{15})$ -;

R₁₁ is -H, halogen, -CN, an aliphatic group, a substituted aliphatic group, an aromatic group, a substituted aromatic group, -OH, -O-(aliphatic group), -O-(substituted aliphatic group), -S-(aliphatic groups), -S-(substituted aliphatic groups), -NO₂, -NH₂, -NH(aliphatic group), -NH(substituted aliphatic group), -N(aliphatic group)₂, -N(substituted aliphatic group)₂;

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 R_{12} an aromatic group or an aliphatic group;

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each R_{13} is independently chosen and is -H, an aliphatic group or a substituted aliphatic group;

n is an integer from one to about four;

B is $-N(R_{16})$ -, -S-, -O- or a covalent bond; and

 R_{14} , R_{15} and R_{16} are independently an aliphatic or substituted aliphatic group, and can be the same or different;

Q is an aliphatic group, a substituted aliphatic group, a benzylic group, a substituted benzylic group, an aromatic group, a substituted aromatic group, a non-aromatic heterocyclic group or a substituted nonaromatic heterocyclic group; and

wherein B, Q and the terminal olefin carbon, taken together, can form a substituted or unsusbstituted non-aromatic heterocyclic ring which is optionally fused to a substituted or unsubstituted aromatic group.

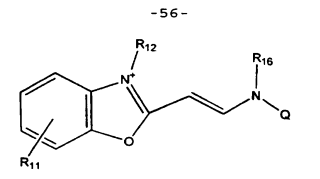
32. The method of Claim 31 wherein:

n is 1;

B is -N(R₁₆)-, -S-, -O- or a covalent bond; and Q is an aliphatic group, a substituted aliphatic group, a benzylic group, a substituted benzylic group, an aromatic group, a substituted aromatic group, a non-aromatic heterocyclic group or a substituted nonaromatic heterocyclic group.

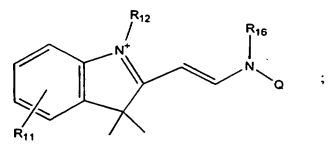
33. The method of Claim 31 wherein the compound is represented by the following structural formula:

BNSDOCID: <WO___9802151A2_I_>



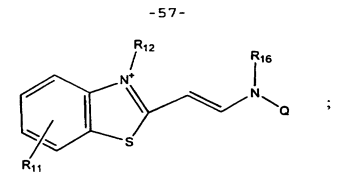
and physiologically acceptable salts thereof.

34. The method of Claim 31 wherein the compound is represented by the following structural formula:



and physiologically acceptable salts thereof.

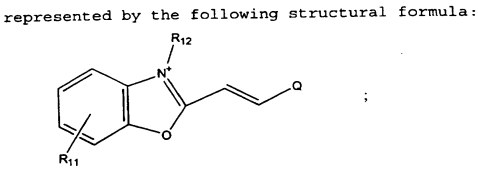
35. The method of Claim 31 wherein the compound is represented by the following structural formula:



and physiologically acceptable salts thereof.

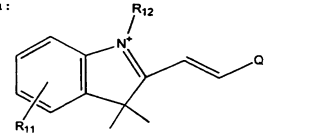
36. The method of Claim 31 wherein the compound is

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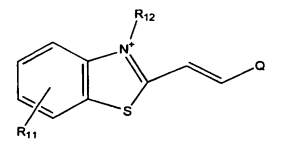
and physiologically acceptable salts thereof.

37. The method of Claim 31 wherein the compound is represented by the following structural formula:
B:2



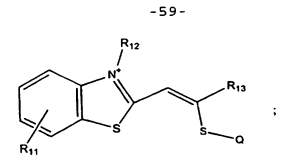
and physiologically acceptable salts thereof.

5 38. The method of Claim 31 wherein the compound is represented by the following structural formula:



and physiologically acceptable salts thereof.

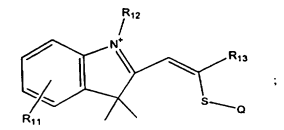
39. The method of Claim 31 wherein the compound is represented by the following structural formula:



and physiologically acceptable salts thereof.

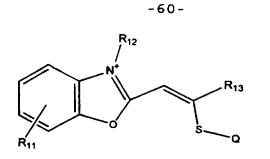
40. The method of Claim 31 wherein the compound is represented by the following structural formula:

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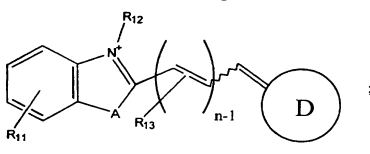
and physiologically acceptable salts thereof.

41. The method of Claim 31 wherein the compound is represented by the following structural formula:



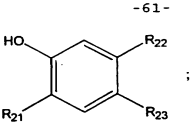
and physiologically acceptable salts thereof.

42. The method of Claim 31 wherein the compound is represented by the following structural Formula:



wherein Ring D is a substituted or unsubstituted non-aromatic heterocyclic ring which is optionally fused to a substituted or unsubstituted aromatic group.

10 43. A method of treating a subject with a disease associated with aberrant leukocyte recruitment and/or activation comprising administering to the individual a therapeutically effective amount of a compound represented by the following structural formula:



and physiologically acceptable salts thereof, wherein:

R₂₁ is -OH, an aliphatic group, a substituted aliphatic group, -O-(aliphatic group), -O-(substituted aliphatic group), -O-CO-(aliphatic group) or -O-CO-(substituted aliphatic group;

 R_{22} and R_{23} are independently -H, an aliphatic group, a substituted aliphatic group, an aromatic group, a substituted aromatic group, -S-(aliphatic group), -S-(substituted aliphatic group), -O-(aliphatic group), -O-(substituted aliphatic group), -(CH₂)_n-R₂₆, and, taken together, can be a -(CH₂)₂- to -(CH₂)₅- alkylene group or a -(CH₂)₂- to -(CH₂)₅- alkylene group substituted with one or more aliphatic groups, substituted aliphatic groups, aromatic groups or substituted aromatic groups; and

 R_{26} is a substituted or unsubstituted aromatic group.

44. The method of Claim 43 wherein:

 R_{21} is -OH, an alkyl group, an alkoxy group, an acetoxy group or an alkyl group substituted with $-NR_{24}R_{25}$;

 R_{22} and R_{23} are independently an alkyl group, an aromatic group, an aralkyl group, and ethylene- R_{26} or thioalkyl, and, taken together, form an alkylene group;

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 R_{24} and R_{25} are independently an alkyl group, an aralkyl group and an aryl group;

 R_{26} is a phenyl group substituted by R_{27} and $R_{28};$ and

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 R_{27} and R_{28} are independently -H, -OH, alkoxy, or halogen.

45. The method of Claim 43 wherein:

 R_{21} is -OH, CH_3CO-O- or an alkyl group substituted with CH_3NH-;

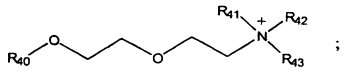
 R_{22} is thioalkyl, alkyl or phenyl; and

 R_{23} is -H, methyl or, taken together with R_{22} , a propylene group, wherein the propylene group is unsubstituted or substituted with one or more methyl or ethyl groups.

46. The method of Claim 44 wherein:

 R_{21} is -OH, CH₃CO-O- or -CH(-CH(CH₃)₂)(-CH₂NHCH₃); R_{22} is -SC₇H₁₅, methyl or phenyl; and R_{23} is -H, methyl or, taken together with R_{22} , a -CH₂CH₂C(CH₃)₂- group.

47. A method of treating a subject with a disease associated with aberrant leukocyte recruitment and/or activation comprising administering to the individual a therapeutically effective amount of a compound represented by the following structural formula:



and physiologically acceptable salts thereof, wherein:

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 R_{40} and R_{43} are independently an aliphatic group, a substituted aliphatic group, a benzylic group, a substituted benzylic group, an aromatic group, a substituted aromatic group, a non-aromatic heterocyclic group or a substituted non-aromatic heterocyclic group;

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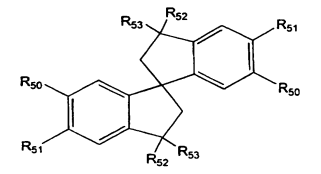
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 R_{41} and R_{42} are independently an aliphatic group or a substituted aliphatic group.

48. The method of Claim 47 wherein R_{41} and R_{42} are each a methyl group.

49. A method of treating a subject with a disease associated with aberrant leukocyte recruitment and/or activation comprising administering to the individual a therapeutically effective amount of a compound represented by the following structural formula:



wherein:

 R_{50} and R_{51} are independently -OH, a halogen, -O-(aliphatic group), -O-(substituted aliphatic group), -O-CO-(aliphatic group), -O-CO-(substituted aliphatic group), -NH₂, -NH(aliphatic group), -NH(substituted aliphatic group), $-N(aliphatic group)_2$, -N(substituted)

aliphatic group)₂, -S-(aliphatic group) or -S-(substituted aliphatic group; and

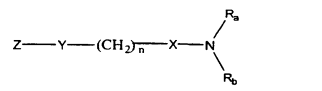
 R_{52} and R_{53} are independently -H, an aliphatic group, a substituted aliphatic group, a halogen, $-NH_2$, -NH(aliphatic group), -NH(substituted aliphaticgroup), $-N(aliphatic group)_2$ or -N(substituted $aliphatic group)_2$.

50. The method of Claim 49 wherein:

 R_{50} and R_{51} are independently -OH, a halogen, -O- (aliphatic group) or -O-(substituted aliphatic group); and

 R_{52} and R_{53} are independently an aliphatic group, a substituted aliphatic group or a halogen.

51. Use of a compound for the manufacture of a medicament for the treatment or prevention of a disease in a subject, said disease being associated with aberrant leukocyte recruitment and/or activation, and said compound being represented by the following structural formula:



and physiologically acceptable salts thereof, wherein:

Z is a substituted or unsubstituted aromatic group;

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Y is a covalent bond, -O- or -CO-; n is an integer from one to about five; X is a covalent bond or -CO-; and

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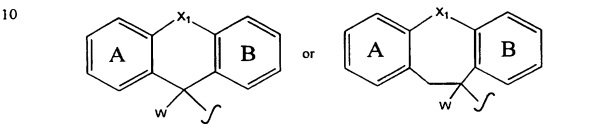
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 R_a is an aliphatic or a substituted aliphatic group; and

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 R_b is an aliphatic group substituted with an aromatic group or substituted aromatic group; and wherein R_a and R_b , taken together with the nitrogen atom bonded to R_a and R_b , can form a substituted or unsubstituted non-aromatic heterocyclic ring.

52. The use of Claim 51 wherein Z is represented by a structural formula selected from:



wherein:

X₁ is a chemical bond, -S-, -CH₂- or -CH₂S-; W is -H, -CN, alkylsulfonyl, carboxamido or

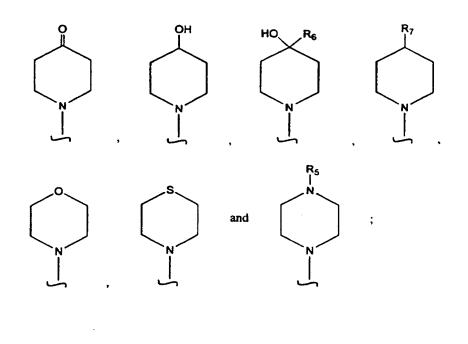
15 carboxyalkyl;

n is an integer from 2-5;

Ring A is substituted with R_{θ} and R_{θ} , wherein R_{θ} and R_{θ} are independently -H, a halogen, alkoxy or alkyl, or, taken together with ring A, form a naphthyl group;

 R_a and R_b are independently -H, alkyl, aralkyl or, taken together with the nitrogen atom bonded to R_a and R_b , form a non-aromatic heterocyclic ring represented by a structure selected from:

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and physiologically acceptable salts thereof, wherein:

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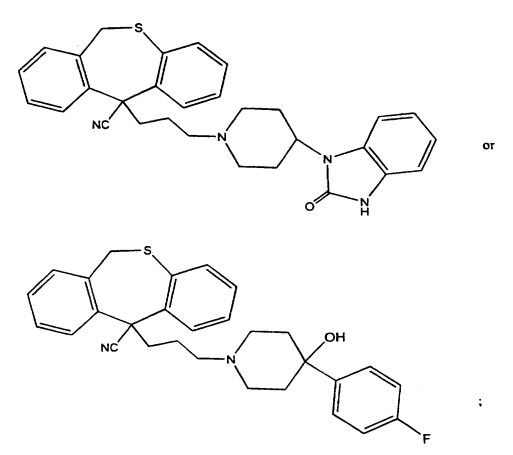
 R_s is -H, alkanoyl, aroyl, aralkoyl, alkyl, aralkyl or cycloalkyl;

 R_6 is an aryl group; and R_7 is -H or a heterocylic ring.

53. The use of Claim 51 wherein the compound is10 represented by a structural formula selected from:

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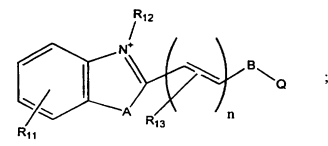


and physiologically acceptable salts thereof.

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54. Use of a compound for the manufacture of a medicament for the treatment or prevention of a disease in a subject, said disease being associated with aberrant leukocyte recruitment and/or activation, and said compound being represented by the following structural formula:



and physiologically acceptable salts thereof, wherein:

A is $>NR_{14}$, $-O_-$, $-S_-$, $-CH_2_-$, $-CH(R_{14})_-$ or $-C(R_{14}R_{15})_-$;

R₁₁ is -H, halogen, -CN, an aliphatic group, a substituted aliphatic group, an aromatic group, a substituted aromatic group, -OH, -O-(aliphatic group), -O-(substituted aliphatic group), -S-(aliphatic groups), -S-(substituted aliphatic groups), -NO₂, -NH₂, -NH(aliphatic group), -NH(substituted aliphatic group), -N(aliphatic group)₂, -N(substituted aliphatic group)₂;

 R_{12} an aromatic group or an aliphatic group; each R_{13} is independently chosen and is -H, an

aliphatic group or a substituted aliphatic group;

n is an integer from one to about four;

B is $-N(R_{16})$ -, -S-, -O- or a covalent bond; and

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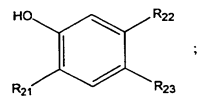
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 R_{14} , R_{15} and R_{16} are independently an aliphatic or substituted aliphatic group, and can be the same or different;

Q is an aliphatic group, a substituted aliphatic group, a benzylic group, a substituted benzylic group, an aromatic group, a substituted aromatic group, a non-aromatic heterocyclic group or a substituted nonaromatic heterocyclic group; and

wherein B, Q and the terminal olefin carbon, 10 taken together, can form a substituted or unsusbstituted non-aromatic heterocyclic ring which is optionally fused to a substituted or unsubstituted aromatic group.

55. Use of a compound for the manufacture of a medicament for the treatment or prevention of a disease in a subject, said disease being associated with aberrant leukocyte recruitment and/or activation, and said compound being represented by the following structural formula:

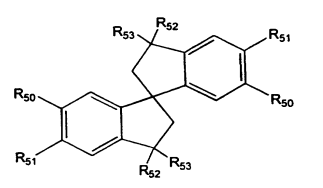


and physiologically acceptable salts thereof, wherein:

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R₂₁ is -OH, an aliphatic group, a substituted aliphatic group, -O-(aliphatic group), -O-(substituted aliphatic group), -O-CO-(aliphatic group) or -O-CO-(substituted aliphatic group;

5		R_{22} and R_{23} are independently -H, an aliphatic group, a substituted aliphatic group, an aromatic group, a substituted aromatic group, -S-(aliphatic group), -S-(substituted aliphatic group), -O-(aliphatic group), -O-(substituted aliphatic group), -(CH_2) _n - R_{26} , and, taken together, can be a -(CH_2) ₂ - to -(CH_2) ₅ - alkylene group or a -(CH_2) ₂ - to -(CH_2) ₅ - alkylene group substituted with one or more aliphatic groups, substituted aliphatic groups,
10		aromatic groups or substituted aromatic groups; and R_{26} is a substituted or unsubstituted aromatic group.
15	56.	The use of Claim 55 wherein: R_{21} is -OH, an alkyl group, an alkoxy group, an acetoxy group or an alkyl group substituted with -NR ₂₄ R ₂₅ ;
20		R_{22} and R_{23} are independently an alkyl group, an aromatic group, an aralkyl group, and ethylene- R_{26} or thioalkyl, and, taken together, form an alkylene
20		<pre>group; R₂₄ and R₂₅ are independently an alkyl group, an aralkyl group and an aryl group; R₂₆ is a phenyl group substituted by R₂₇ and R₂₈;</pre>
25		and R ₂₇ and R ₂₈ are independently -H, -OH, alkoxy, or halogen.
	57.	Use of a compound for the manufacture of a medicament for the treatment or prevention of a disease in a subject, said disease being associated with aberrant
30		leukocyte recruitment and/or activation, and said compound being represented by the following structural formula:



wherein:

 R_{50} and R_{51} are independently -OH, a halogen, -O-(aliphatic group), -O-(substituted aliphatic group), -0-CO-(aliphatic group), -0-CO-(substituted aliphatic group), -NH₂, -NH(aliphatic group), -NH(substituted aliphatic group), -N(aliphatic group)₂, -N(substituted aliphatic group)2, -S-(aliphatic group) or -S-(substituted aliphatic group; and

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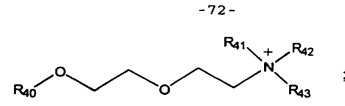
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 R_{52} and R_{53} are independently -H, an aliphatic group, a substituted aliphatic group, a halogen, $-NH_2$, -NH(aliphatic group), -NH(substituted aliphatic group), $-N(aliphatic group)_2$ or -N(substitutedaliphatic group)₂.

58. Use of a compound for the manufacture of a medicament 15 for the treatment or prevention of a disease in a subject, said disease being associated with aberrant leukocyte recruitment and/or activation, and said compound being represented by the following structural formula:

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and physiologically acceptable salts thereof, wherein:

R₄₀ and R₄₃ are independently an aliphatic group, a substituted aliphatic group, a benzylic group, a substituted benzylic group, an aromatic group, a substituted aromatic group, a non-aromatic heterocyclic group or a substituted non-aromatic heterocyclic group;

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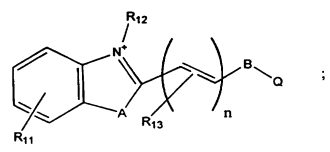
 R_{41} and R_{42} are independently an aliphatic group or a substituted aliphatic group.

59. A pharmaceutical composition comprising the compound of Claim 31 and a suitable pharmaceutical carrier.

60. A pharmaceutical composition comprising the compound 15 of Claim 43 and a suitable pharmaceutical carrier.

- 61. A pharmaceutical composition comprising the compound of Claim 44 and a suitable pharmaceutical carrier.
- 62. A pharmaceutical composition comprising the compound of Claim 47 and a suitable pharmaceutical carrier.
- 20 63. A pharmaceutical composition comprising the compound of Claim 49 and a suitable pharmaceutical carrier.
 - 64. A compound for use in the treatment or prevention of a disease in a subject, said disease being associated with aberrant leukocyte recruitment and/or activation.

and said compound being represented by the following structural formula:



	and physiologically acceptable salts thereof,
5	wherein:
	A is NR_{14} , -O-, -S-, -CH ₂ -, -CH(R_{14}) - or
	$-C(R_{14}R_{15}) - ;$
	R_{11} is -H, halogen, -CN, an aliphatic group, a
	substituted aliphatic group, an aromatic group, a
10	substituted aromatic group, -OH, -O-(aliphatic group),
	-O-(substituted aliphatic group), -S-(aliphatic
	groups), -S-(substituted aliphatic groups), -NO ₂ , -NH ₂ ,
	-NH(aliphatic group), -NH(substituted aliphatic
	group), -N(aliphatic group) ₂ , -N(substituted aliphatic
15	group) ₂ ;
	R_{12} an aromatic group or an aliphatic group;
	each R_{13} is independently chosen and is -H, an
	aliphatic group or a substituted aliphatic group;

n is an integer from one to about four;

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B is $-N(R_{16})$ -, -S-, -O- or a covalent bond; and

 $R_{14},\ R_{15}$ and R_{16} are independently an aliphatic or substituted aliphatic group, and can be the same or different;

Q is an aliphatic group, a substituted aliphatic group, a benzylic group, a substituted benzylic group, an aromatic group, a substituted aromatic group, a non-aromatic heterocyclic group or a substituted nonaromatic heterocyclic group; and

wherein B, Q and the terminal olefin carbon, taken together, can form a substituted or

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unsusbstituted non-aromatic heterocyclic ring which is optionally fused to a substituted or unsubstituted aromatic group.

65. A compound for use in the treatment or prevention of a disease in a subject, said disease being associated

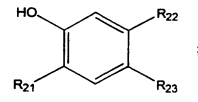
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with aberrant leukocyte recruitment and/or activation, and said compound being represented by the following structural formula:



and physiologically acceptable salts thereof, wherein:

R₂₁ is -OH, an aliphatic group, a substituted aliphatic group, -O-(aliphatic group), -O-(substituted aliphatic group), -O-CO-(aliphatic group) or -O-CO-(substituted aliphatic group;

R₂₂ and R₂₃ are independently -H, an aliphatic group, a substituted aliphatic group, an aromatic group, a substituted aromatic group, -S-(aliphatic group), -S-(substituted aliphatic group),

-O-(aliphatic group), -O-(substituted aliphatic group), -(CH_2)_n-R₂₆, and, taken together, can be a -(CH_2)₂- to -(CH_2)₅- alkylene group or a -(CH_2)₂- to -(CH_2)₅- alkylene group substituted with one or more

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aliphatic groups, substituted aliphatic groups, aromatic groups or substituted aromatic groups; and R_{26} is a substituted or unsubstituted aromatic group.

5 66. The compound of Claim 65 wherein:

 R_{21} is -OH, an alkyl group, an alkoxy group, an acetoxy group or an alkyl group substituted with -NR₂₄R₂₅;

 R_{22} and R_{23} are independently an alkyl group, an aromatic group, an aralkyl group, and ethylene- R_{26} or thioalkyl, and, taken together, form an alkylene group;

 R_{24} and R_{25} are independently an alkyl group, an aralkyl group and an aryl group;

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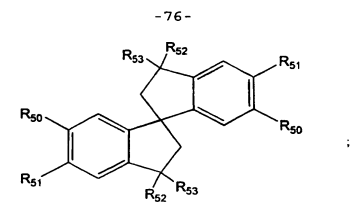
 R_{26} is a phenyl group substituted by R_{27} and $R_{20};$ and

 R_{27} and R_{28} are independently -H, -OH, alkoxy, or halogen.

67. A compound for use in the treatment or prevention of a

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disease in a subject, said disease being associated with aberrant leukocyte recruitment and/or activation, and said compound being represented by the following structural formula:



wherein:

 R_{50} and R_{51} are independently -OH, a halogen, -O-(aliphatic group), -O-(substituted aliphatic group), -O-CO-(aliphatic group), -O-CO-(substituted aliphatic group), -NH₂, -NH(aliphatic group), -NH(substituted aliphatic group), -N(aliphatic group)₂, -N(substituted aliphatic group)₂, -S-(aliphatic group) or -S-(substituted aliphatic group; and

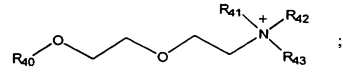
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 R_{52} and R_{53} are independently -H, an aliphatic group, a substituted aliphatic group, a halogen, -NH₂, -NH(aliphatic group), -NH(substituted aliphatic group), -N(aliphatic group)₂ or -N(substituted aliphatic group)₂.

15 68. A compound for use in the treatment or prevention of a disease in a subject, said disease being associated with aberrant leukocyte recruitment and/or activation, and said compound being represented by the following structural formula:

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and physiologically acceptable salts thereof, wherein:

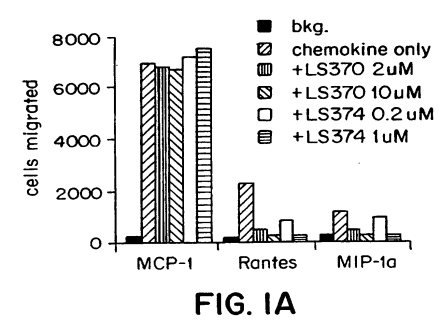
 R_{40} and R_{43} are independently an aliphatic group, a substituted aliphatic group, a benzylic group, a substituted benzylic group, an aromatic group, a substituted aromatic group, a non-aromatic heterocyclic group or a substituted non-aromatic heterocyclic group;

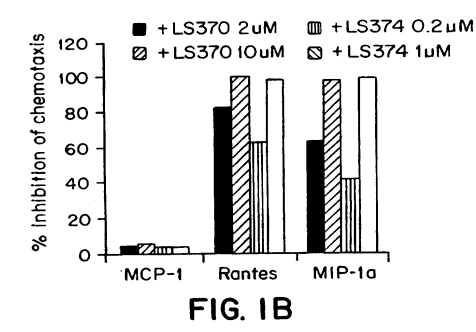
 R_{41} and R_{42} are independently an aliphatic group or a substituted aliphatic group.

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INTERNATIONAL APPLICATION PUBLIS		
(51) International Patent Classificati n 6 :		(11) International Publication Number: WO 98/02151
A61K 31/13, 31/135, 31/445, 31/495, 31/535, 31/54, 31/38	A3	(43) International Publication Date: 22 January 1998 (22.01.98)
 (21) International Application Number: PCT/US4 (22) International Filing Date: 11 July 1997 (13) (30) Priority Data: 60/021,716 12 July 1996 (12.07.96) (71) Applicant: LEUKOSITE, INC. [US/US]; 215 Fire Cambridge, MA 02142 (US). (72) Inventors: SCHWENDER, Charles, F.; 577 East H Glen Gardner, NJ 08826 (US). MACKAY, Charles Church Street, Watertown, MA 02172 (US). Julia, C.; 8 Chubb's Brook Lane, Beverly Farms, M (US). NEWMAN, Walter; 8 Durham Street No. 3 MA 02115 (US). (74) Agents: BROOK, David, E. et al.; Hamilton, Brook, Reynolds, P.C., Two Militia Drive, Lexington, M (US). 	11.07.9 U st Stre Iill Roa arles, I PINT IA 019 3, Bosto Smith	 BY, CA, CH, CN, CU, CZ, DE, DK, EE, ES, FI, GB, GE, GH, HU, IL, IS, JP, KE, KG, KP, KR, KZ, LC, LK, LR, LS, LT, LU, LV, MD, MG, MK, MN, MW, MX, NO, NZ, PL, PT, RO, RU, SD, SE, SG, SI, SK, SL, TJ, TM, TR, TT, UA, UG, UZ, VN, YU, ZW, ARIPO patent (GH, KE, LS, MW, SD, SZ, UG, ZW), Eurasian patent (AM, AZ, BY, KG, KZ, MD, RU, TJ, TM), European patent (AT, BE, CH, DE, DK, ES, FI, FR, GB, GR, IE, IT, LU, MC, NL, PT, SE), OAPI patent (BF, BJ, CF, CG, CI, CM, GA, GN, ML, MR, NE, SN, TD, TG). Published With international search report. Before the expiration of the time limit for amending the claims and to be republished in the event of the receipt of amendments. & (88) Date of publication of the international search report:
integer from one to about five X is a covalent bond or -	d or ur CO R c grou	$-(CH_2)_n - X - N_R_b$ (1) asubstituted aromatic group. Y is a covalent bond, -O- or -CO n is an a is an aliphatic or a substituted aliphatic group; R_b is an aliphatic group; R_b is an aliphatic group; R_b is an aliphatic group; R_b and R_b , can be considered as the second

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INTERNATION SEARCH

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A. CLASSIFICATION OF SUBJECT MATTER

A 61 K 31/13,A 61 K 31/135,A 61 K 31/445,A 61 K 31/495, A 61 K 31/535,A 61 K 31/54,A 61 K 31/38

According to International Patent Classification (IPC) or to both national classification and IPC 6

B. FIELDS SEARCHED

Minimum documentation searched (classification system followed by classification symbols)

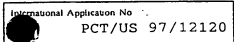
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Documentation searched other than minimum documentation to the extent that such documents are included in the fields searched

Electronic data base consulted during the international search (name of data base and, where practical, search terms used)

C. DOCUMENTS CONSIDERED TO BE RELEVANT Relevant to claim No. Citation of document, with indication, where appropriate, of the relevant passages Category * 1-3,6, HELWIG, H. et al. Arznei-Х 18,19, mittel. Stuttgart: Helwig/ Otto Arzneimittel, 1992. 51 Vol. 1, 8th edition, pages 4-1 to 4-24, especially 4-8: Thiethylparazin. 1,2,6, Chem. abstr., Vol. 104, Α No. 5, 03 February 1986 18-29, 51-53 (Columbus, Ohio, USA), page 540, column 2, the abstract No. 33990s, SINDELAR, K. et al. "Potential antidiarrheal agents: 1-(11-cyano-6,11-dihydrodibenzo(b,e)thiepin-11-ylalkyl)-and 1--(10-cyano-10,11-dihydrodibenzo(b,f)thiepin-10-ylalkyl)-4-substituted piperi-Patent family members are listed in annex. Further documents are listed in the continuation of box C. x I * Special categories of cited documents : "T" later document published after the international filing date or priority date and not in conflict with the application but "A" document defining the general state of the art which is not ... cited to understand the principle or theory underlying the considered to be of particular relevance invention .Е. earlier document but published on or after the international "X" document of particular relevance; the claimed invention cannot be considered novel or cannot be considered to filing date involve an inventive step when the document is taken alone "L" document which may throw doubts on priority claim(s) or which is cited to establish the publication date of another citation or other special reason (as specified) "Y" document of particular relevance; the claimed invention cannot be considered to involve an inventive step when the document is combined with one or more other such docu-.0. document referring to an oral disclosure, use, exhibition or ments, such combination being obvious to a person skilled other means in the art. document published prior to the international filing date but later than the pnority date claimed .Ь. "&" document member of the same patent family Date of mailing of the international search report Date of the actual completion of the international search 03 November 1997 0 4. 05. 98 Authorized officer Name and mailing address of the ISA European Patent Office, P.B. 5818 Patentiaan 2 NL - 2280 HV Rijswijk Tel. (+ 31-70) 340-2040, T'x. 31 651 epo nl, MAZZUCCO e.h. Fax: (+31-70) 340-3016 Form PCT/ISA/210 (second sheet) (July 1992)

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C.(Conunua	uon) DOCUMENTS CONSIDERED TO BE RELEVANT	
Category *	Citation of document, with indication, where appropriate, of the relevant passages	Reievant to claim No.
A	<pre>dines," Collect. Czech. Chem. Commun. 1985, 50(5), 1089-96 (Eng) (cited in the application). Chem. abstr., Vol. 109, No. 11, 12 September 1988 (Columbus, Ohio, USA), page 689, column 2, the abstract No. 92794g, PROTIVA, M. et al. "Substituted 11-(pi- peridinoalkyl)-6,11-dihy- drodibenzo(b,e)thiepin-11- -carbonitriles useful as antidiarrheal drugs," Czech. CS 240,698</pre>	1,2,6, 18-23, 27-29, 51-53
A	<pre>(cited in the application). WO 90/13539 A1 (MEIJI SEIKA KAISHA, LTD.) 15 November 1990 (15.11.90), abstract (cited in the application).</pre>	1-3,9- 10,51
A	US 4086234 A (DRYDEN, H.L. et al.) 25 April 1978 (25.04.78), abstract, claims 1,4,5, examples 2-4 (cited in the application).	1,2,6, 16,17, 51
A	US 3922266 A (KATSUBE, J. et al.) 25 November 1975 (25.11.75), abstract, column 1, line 1 - column 2, line 23, column 3, lines 14-18 (cited in the application).	1,2,3, 6,14- 17,51
А	US 3936468 A (YAMAMOTO, H. et al.) 03 February 1976 (03.02.76), abstract, column 1, lines 6-60 (cited in the application).	1,2,6, 16,17; 51
A	US 3907812 A (YAMAMOTO, H. et al.) 23 September 1975 (23.09.75), abstract, column 1, lines 15-67 (cited in the application).	1,2,6, 16,17, 51

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	INTERNATIONAL SEARCH REP	PORT						
<u> </u>			PCT/US 97/ 12120					
Box i	Observations where certain claim were fund	i unsearchabl (Continu	uation of it in 1 of first sheet)					
This Inte	s International Search Report has not been established in respect of certain claims under Article 17(2)(a) for the following reasons:							
1. X 2.	Although claims 1-50 are directed to a method of treatment of the human or the animal body by therapy, the search has been carried out for the matter of claims 1-30 and has been based on the alleged effects of the composition (see PCT Rule 39.1 (iv)).							
^{د.} لـــا	because they relate to parts of the International Applica an extent that no meaningful International Search can t	tion that do not comply with t e carried out, specifically:	the prescribed requirements to such					
3.	Claims Nos.: because they are dependent claims and are not drafted	l in accordance with the seco	ond and third sentences of Rule 6.4(a).					
Box il	Observations where unity of invention is lack	ing (Continuation of iter	n 2 of first sheet)					
This Int	ernational Searching Authority found multiple inventions	in this international application	on, as follows:					
ระ ระ ระ	ubject 1: 1-30, 51-53 ubject 2: 31-42, 54, 59, 64 ubject 3: 43-46, 55, 56, 60, 61, 6 ubject 4: 47, 48, 58, 62, 68 ubject 5: 49-50, 57, 63, 67	5, 66						
	bjeet 5. 45 56, 67, 66, 67							
1.	As all required additional search fees were timely paid searchable claims.	by the applicant, this Interna	tional Search Report covers all					
1	As all required additional search fees were timely paid							
	As all required additional search fees were timely paid searchable claims. As all searchable claims could be searched without eff	ort justifying an additional fee ere timely paid by the applica	e, this Authority did not invite payment					
2.	As all required additional search fees were timely paid searchable claims. As all searchable claims could be searched without eff of any additional fee. As only some of the required additional search fees we covers only those claims for which fees were paid, spe	ort justifying an additional fee ere timely paid by the applica cifically claims Nos.:	e, this Authority did not invite payment nt, this International Search Report					

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ANHANG

- zum internationalen Recherchen-bericht über die internationale Patentanmeldung Nr.

ANNEX

to the International Search Report to the International Patent Application No.

FCT/US 97/12120 SAE 166629

In diesee Anhang sind die Mitglieder der Patentfamilien der im obenge-nannten internationalen Recherchenbericht Diese Angaben dienen nur zur Unter-richtung und erfolgen ohne Gewähr. This Annex lists the patent family members relating to the patent documents national search report. The Office is in no way liable for these particulars which are given merely for the purpose of information.

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ANNEXE

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La presénte annexe indique les membres de la famille de brevets relatifs aux documents de brevets cités dans le rapport de recherche inter-national visée ci-dessus. Les reseigne-ments fournis sont donnés à titre indica-tif et n'engagent pas la responsibilité de l'Office.

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angeführtes Patent o in sea Document o	erchenbericht 5 Patentdokuæent document cited rch report de brevet cité oport de recherche	Datum der Veröffentlichung Publication date Date de publication	Mitglied(er) der Patentfamilie Patent family member(s) Membre(s) de la famille de brevets	Datum der Veröffentlichung Publication date Date de publication				
WO A1	9013539	15-11-90	keine – none –)	rien				
US A	4086234	25-04-78	AU A1 19297/76 AU A1 19297/76 AU A1 1064925 CCE A1 062509661 FRE A1 2655096677 GBP A2 2655006677 GBP A2 2655006677 GBP A2 26550096477 GBP A2 2655009742284 JPLLE A 76123779 JPLLE A 76128469 SEE C 428469	$\begin{array}{c} 11 - 05 - 78\\ 211 - 02 - 80\\ 233 - 101 - 81\\ 155 - 05 - 77\\ 055 - 05 - 77\\ 055 - 064 - 77\\ 155 - 064 - 87\\ 156 - 055 - 87\\ 156 - 055 - 87\\ 268 - 055 - 87\\ 065 - 055 - 77\\ 084 - 07 - 83\\ 13 - 10 - 83\end{array}$	- 24 ș			
US A	3922266	25-11-75	649 640 640 640 640 640 640 640 640 640 640 640 640	$\begin{array}{c} 183 \\ 183 \\ -777 \\ 776 \\ -777 $				
	3936468	03-02-76	ATE AA1 9789401 057994 057994 057994 05799401 0799401 0799400 07743050200 07743050200 051050200 05105000 0000 00000 00000 00000 00000 00000 0000	$\begin{array}{c} 11 - 12 - 72 \\ 99 - 69 - 69 - 75 \\ 03 - 02 - 76 \\ 31 - 02 - 74 \\ 27 - 12 - 74 \\ 02 - 09 - 75 \\ 03 - 00 - 71 \\ 18 - 09 - 75 \\ 28 - 09 - 95 \\ 28 - 09 - 95 \\ 28 - 09 - 95 \\ 28 - 09 - 95 \\ 28 - 09 - 95 \\ 28 - 09 - 95 \\ 28 - 09 - 95 \\ 28 - 09 - 95 \\ 28 - 09 - 95 \\ 28 - 09 - 95 \\ 28 - 09 - 95 \\ 28 - 09 - 95 \\ 28 - 09 - 95 \\ 28 - 09 - 95 \\ 28 - 09 - 95 \\ 28 - 09 - 95 \\ 28 - 09 - 95 \\ 28 - 09 - 95 \\ 28 - 09 - 95 \\ 28 - 09 \\ 28 - 09 - 95 \\ 28 - 09 \\ 28 -$				

•			7777333440004300430043004300430043004300	$\begin{array}{c} 10 - 11 - 75 \\ 30 - 11 - 788 \\ 10 - 778 \\ 10 - 7$
			P 104257 SE B 396383 SE C 396383 JP B4 50004672 SU D 421192 FR A1 2070146 FR B1 20701466 FR B1 20701466 ZA A 7007192	31-08-79 19-09-77 29-12-77 29-12-75 25-03-74 10-09-71 10-09-71 22-02-74 28-07-71
US A	3907812	23-09-75	keine – none – r	ien

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